

REMARKS

Examiner J. L. Brophy is thanked for the thorough examination and search of the subject Patent Application. Claims 5 and 16 have been amended.

All Claims are believed to be in condition for Allowance and that is so requested.

We apologize for neglecting to respond to the Double patenting rejection. Claims 5 and 16 have been amended to include the roughened surface of the silicon-implanted low-k dielectric layer taught on the top of page 8 of the Specification. It is believed that this amendment overcomes the double patenting objection to these claims over Claims 24 and 28.

Reconsideration of the rejection under 35 U.S.C. 102 of Claims 1, 3 and 6 as being anticipated by JP08102489A is requested in accordance with the following remarks.

Applicants' invention provides a means for improving adhesion of a TEOS oxide layer to an underlying low dielectric constant material layer. Silicon ions are

implanted into the underlying layer to improve adhesion. As shown in Fig. 2, the silicon implantation is performed on a flat surface. The underlying passivation layer 12 is also a planarizing layer. In JP08102489A, the silicon implantation is performed over non-planar features (See Fig. 1a).

JP08102489A teaches implanting silicon ions into a plasma TEOS film to prevent moisture diffusion. Those skilled in the art of microelectronics know that the definition of low dielectric constant means a dielectric constant of less than 3. (See Exhibits A and B). The dielectric constants of the group of low dielectric constant materials in Applicants' invention: i.e., porous or non-porous carbon-based silicon oxides, porous or non-porous doped silicon oxides, porous or non-porous organic polymers, or porous or non-porous inorganic polymers are all less than 3. (See Exhibit C for doped silicon oxides, Exhibit D for carbon-based silicon oxides, and Exhibit E for organic and inorganic polymers) The dielectric constant of PECVD TEOS is 4 to 4.2 (see Exhibit F). Thus, the material of JP08102489A is not a low dielectric constant material and is therefore different from Applicants' claimed invention.

Reconsideration of the rejection under 35 U.S.C. 102 of Claims 1, 3 and 6 as being anticipated by JP08102489A is requested in accordance with the remarks above.

Reconsideration of the rejection under 35 U.S.C. 102 of Claim 1 as being anticipated by Muroyama is requested in accordance with the following remarks.

Muroyama teaches implanting silicon ions into a thermal oxide film to make its surface hydrophobic. The dielectric constant of undoped silicon dioxide is 4 to 4.2. (See Exhibit F) Thus, the material of Muroyama is not a low dielectric constant material and is therefore different from Applicants' claimed invention.

Reconsideration of the rejection under 35 U.S.C. 102 of Claim 1 as being anticipated by Muroyama is requested in accordance with the remarks above.

Reconsideration of the rejection under 35 U.S.C. 102 of Claims 1, 3 and 6 as being anticipated by Wantanabe et al is requested in accordance with the following remarks.

Watanabe et al teaches implanting silicon ions into a spin-on-glass film to decompose its organic components. Organic spin-on-glass appears to fall into the category of low dielectric constant materials of carbon-doped silicon oxide. However, the precursor of $(\text{CH}_3\text{Si}(\text{OH})_3)$ (col. 6, lines

31-36) will lead to a final product of carbon-based silicon dioxide plus carbon-doped silanol (with Si-OH bonds). This is well-known for SOG. Furthermore, those skilled in the art of microelectronics will know that the dielectric constant of organic SOG is greater than 3 (see Exhibit G). We apologize for neglecting to refer here to Exhibit G in the previous response. Exhibit G was included in the previous response, but not referenced. Thus, the material of Wantanabe et al is not a low dielectric constant material and is therefore different from Applicants' claimed invention.

At issue is the meaning of the term "low dielectric constant", specifically as it relates to spin-on-glass (SOG) materials. Traditional SOG's have been used in the industry since the late 1980's. On pages 419 and 430 of ULSI Technology (Exhibit H), there are basically two types of SOG: organic and inorganic whose structures are shown in Fig. 28. The organic SOG is also known as siloxane SOG and is the one that Wantanabe et al is using. It is important to note the following:

1) Based on the structures, both types of SOG will behave like silicon dioxide. Those skilled in the art know that the term "glass" refers to silicon dioxide. Other examples include PSG (phosphosilicate glass) which is phosphorus doped silicon dioxide. Hence, the term "spin-on-

glass" refers to silicon dioxide that is deposited by spin coating.

2) Organic SOG or siloxane SOG contains carbon and in the third paragraph on page 420 (Exhibit H) "retains carbon even after etching."

3) In the middle portion of the first paragraph on page 421, the dielectric constant of SOG is affirmed. "Although some fully cured SOG's have lower dielectric constants (3.0 to 3.6 compared with 3.9 to 4.5 for SiO₂) than oxide..."

Hence, siloxane SOG has a dielectric constant of more than 3.

On page 508 of the Proceedings from the Symposium on Polymers for Microelectronics, the 1997 "National Technology Roadmap for Semiconductors" (Exhibit I) mentions that the dielectric constant of low k materials is less than 3. Wantanabe's patent was filed on 21 October 1997, and thus reads onto this roadmap. Also, it means that the traditional SOG, silicate or siloxane, has a dielectric constant of greater than 3.

Organic siloxane SOG has a dielectric constant of greater than 3. However, sometime in 1996, there is another spin-on material known as alkyl silsesquioxane that came on

the market. The difference in structure between a siloxane and a silsesquioxane is given in Figure 34 of Stanley Wolf's Contract Course for Chartered (Exhibit J). It can be seen that the silsesquioxane is more structured in the sense that each silicon atom has one alkyl group (denoted by R) attached to it and hence a stoichiometric formula can be ascribed to the silsesquioxane (in this case it is $\text{RSiO}_{1.5}$). It is because of this structural arrangement that the dielectric constant of alkyl silsesquioxane is able to come below 3.0. This is the "non-porous carbon-doped silicon oxide" in Applicants' invention. On the bottom of page 6 of the Specification, alkyl silsesquioxane is listed as an example of this type of material. On the other hand, Fig. 34 shows the alkyl groups to be randomly distributed in the silicon dioxide matrix of the siloxane SOG's. Overall, each silicon atom will have less than one alkyl group attached to it.

Furthermore, there is the presence of Si-OH (silanol groups) on the siloxane SOG's. It is because of these silanol groups that the siloxane and silicate SOG's are rendered hygroscopic; i.e. moisture (water) can attach to the OH groups through hydrogen bonding. The hygroscopic behavior of siloxane SOG's significantly differentiates them from the silsesquioxane (low k SOG) since the latter has significantly higher amounts of carbon and negligible silanol

groups and thus will contain negligible moisture.

Based on Fig. 34 and the above explanations, it is not wrong to classify silsesquioxanes as SOG. If you look at the Table of Low k Dielectric Materials (page 443 of Exhibit I), you will find "spin-on-glasses" as one of the materials. To a layman, this causes confusion. But those skilled in the art will understand without any ambiguity that the spin-on-glasses listed under low k dielectric materials or listed under spin-on dielectrics but with a dielectric constant less than 3 are actually alkyl silsesquioxanes (and not siloxanes). In other words, the confusion will not happen so long as silsesquioxanes are listed as "low k SOG", "low k organic SOG" or simply "silsesquioxanes".

In the Advisory Action, the Examiner refers to Table 1 of Exhibit F where spin on dielectrics (SOD) are shown. The differentiation between spin on dielectrics (SOD) and spin-on-glass (SOG) can be found in Wolf & Tauber (Exhibit K). On page 792, SOG falls under the category of "First Generation Low k Dielectrics ($2.8 < k < 3.5$)" whereas spin on dielectrics (SOD) fall under the category of "Second-Generation Low k Dielectrics ($2.5 < k < 2.8$)" on page 793. As the term spin-on-dielectrics implies, it means any dielectric material that is spun on. SOD refers to non-SOG

materials such as organic polymers as well as low dielectric constant SOG (i.e. silsesquioxane, etc.). In other words, SOG is a subset of SOD, but SOD is not a subset of SOG. The exception to this statement is if the SOD is silsesquioxane.

Watanabe et al states in col. 4, lines 64-67 that the primary objective is "to provide a semiconductor device with less water and hydroxyl groups contained in the SOG." In the Background of the Invention section, he refers to two articles which describe the presence of water in organic SOG's. In the portion of the first article attached as Exhibit L, on page 102, the organic SOG's used are Accuglass 111 and 314. From the AlliedSignal Spin-on Products chart (Exhibit M), these are siloxane SOG (dielectric constant > 3). These cured SOG's have moisture absorption. The second article portion attached as Exhibit N shows that the organic SOG used is siloxane based. Since Watanabe et al referred to these siloxane SOG's in their patent, the organic SOG used in their patent is siloxane SOG since they share the same issue of moisture retention.

As explained above, it is because of these silanol groups that the siloxane SOG's are rendered hygroscopic; i.e. moisture (water) can attach to the OH groups through hydrogen

bonding. In col. 9, lines 47-55, it says "...it can be understood that the water and hydroxyl groups contained in the modified SOG film 7 are reduced immediately after the ion implantation compared with the organic SOG pre film 6." In other words, the organic SOG used by Wantanabe et al contains silanol groups and thus cannot be silsesquioxane or low-k SOG, but will be a siloxane. This is because silsesquioxane has significantly higher amounts of carbon (more hydrophobic) and thus will contain negligible moisture.

It is hoped that the above explanation will show that although the definition of "low dielectric constant" is not found in the Specification, it is well known in the art. Furthermore, it is hoped that the above explanation shows clearly that the low-k materials recited in Applicants' claims are not the same as the siloxane SOG taught by Wantanabe et al.

Reconsideration of the rejection under 35 U.S.C. 102 of Claims 1, 3 and 6 as being anticipated by Wantanabe et al is requested in accordance with the remarks above.

Reconsideration of the rejection under 35 U.S.C. 103 of Claims 4, 5, and 24-27 as being unpatentable over JP08102489A or Wantanabe et al is requested in accordance with the

following remarks.

JP08102489A teaches implanting silicon ions into a plasma TEOS film to prevent moisture diffusion. Wantanabe et al teaches implanting silicon ions into a spin-on-glass film to decompose its organic components. As discussed above, low dielectric constant (<3) materials claimed in Applicants' invention are not taught or suggested in the references.

Reconsideration of the rejection under 35 U.S.C. 103 of Claims 4, 5, and 24-27 as being unpatentable over JP08102489A or Wantanabe et al is requested in accordance with the remarks above.

Reconsideration of the rejection under 35 U.S.C. 103 of Claims 3-6 and 24-27 as being unpatentable over Muroyama is requested in accordance with the following remarks.

Muroyama teaches implanting silicon ions into a thermal oxide film to make its surface hydrophobic. As discussed above, low dielectric constant (<3) materials claimed in Applicants' invention are not taught or suggested in the reference.

Reconsideration of the rejection under 35 U.S.C. 103 of Claims 3-6 and 24-27 as being unpatentable over Muroyama is

requested in accordance with the remarks above.

Reconsideration of the rejection under 35 U.S.C. 103 of Claims 12, 14-17, 23, and 28-32 as being unpatentable over Wantanabe et al is requested in accordance with the following remarks.

Wantanabe et al teaches implanting silicon ions into a spin-on-glass film to decompose its organic components. As discussed above, low dielectric constant (<3) materials claimed in Applicants' invention are not taught or suggested in the references. It is agreed that forming a copper layer within an opening is taught in the reference, but this is not a damascene process.

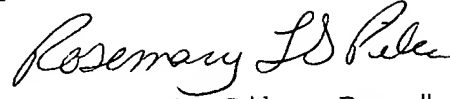
Reconsideration of the rejection under 35 U.S.C. 103 of Claims 12, 14-17, 23, and 28-32 as being unpatentable over Wantanabe et al is requested in accordance with the remarks above.

Allowance of all Claims is requested.

It is requested that should Examiner Brophy not find that the Claims are now Allowable that she call the undersigned at 765 4530866 to overcome any problems preventing allowance.

CS-01-001

Respectfully submitted,

A handwritten signature in cursive script that reads "Rosemary L. S. Pike". The signature is written in dark ink and is positioned above the printed name.

Rosemary L. S. Pike. Reg # 39,332

Attachments: Exhibits A-N

VERSION WITH MARKINGS TO SHOW CHANGES MADE

IN THE CLAIMS

Please amend the Claims as follows:

5. (AMENDED) The method according to Claim 1 wherein said silicon ions are implanted into said low dielectric constant material layer to a depth of between about 50 and 600 Angstroms thereby forming a roughened silicon-implanted surface of said low dielectric constant material layer.

16. (AMENDED) The method according to Claim 12 wherein said silicon ions are implanted into said first and second low dielectric constant material layers to a depth of between about 50 and 600 Angstroms thereby forming a roughened silicon-implanted surface of said first and second low dielectric constant material layers.



LOW DIELECTRIC CONSTANT OXIDE FILMS DEPOSITED USING CVD TECHNIQUES

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ABSTRACT

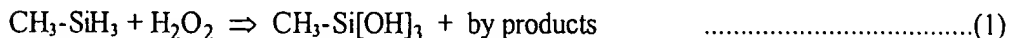
It has been shown previously that low dielectric constant insulating layers can be deposited using a process technology known as "Low-k Flowfill®". This process uses a chemical vapour deposition (CVD) reaction between methyl-silane ($\text{CH}_3\text{-SiH}_3$) and hydrogen peroxide (H_2O_2) to form a methyl doped silicon oxide. Layers deposited using this technique exhibit a dielectric constant < 3.0 and remain stable upon annealing at temperatures of up to 500°C . The layers also exhibit excellent gap-filling capabilities, being able to completely fill 0.1 micron features with up to 4 to 1 aspect ratios. This paper considers the Low-k Flowfill® process in more detail, and examines some of the integration issues that the use of this material raises. The possibility of enhancing the basic methyl-silane/hydrogen peroxide chemistry in order to achieve a lower dielectric constant is also considered. This approach is contrasted with the possibility of using new alternative chemistries in order to achieve dielectric constants values of < 2.5 . The use of the pre-cursors dimethyl-silane and methylenebis-silane are considered.

INTRODUCTION

Flowfill® technology, described elsewhere [1-5], utilises a reaction between the simple inorganic precursors silane (SiH_4) and hydrogen peroxide (H_2O_2) to form a flowable SiO_2 that has the ability to fill deep sub-micron features whilst at the same time providing excellent local and a degree of global planarization. The dielectric constant of this material is in the range 3.9 - 4.4 depending on application. In contrast, the low-k Flowfill® process [6] achieves a dielectric constant of ~ 2.9 whilst maintaining many of the attractive properties of the original Flowfill® process.

LOW-K FLOWFILL® MECHANISM

The key to the low-k Flowfill® process is the reaction between the gaseous pre-cursor methyl silane ($\text{CH}_3\text{-SiH}_3$) and hydrogen peroxide (H_2O_2). Under the correct process conditions these precursor materials react at the surface of the substrate to form what is believed to be a methyl doped silanol type structure. The reaction can be represented in its simplest form by :



Theoretically the CH_3 group does not actively take part in the reaction with the H_2O_2 and as a result remains bonded to the silicon atom. It is postulated that the silanol is then converted into an amorphous methyl doped silicon oxide structure via condensation reactions. This can be represented by:



where $x = 1.5 - 2.0$.

These process steps are depicted in Figure 1. The incorporation of the CH_3 group bound to the silicon atom in the oxide lattice gives rise to the reduction in dielectric constant.

Since the low-k Flowfill[®] material is formed from gaseous pre-cursors, seamless gap-fill down to $0.1\mu\text{m}$, with up to 4:1 aspect ratios can be achieved. The film effectively grows from the bottom-up. At a substrate temperature of 0°C the rate of polymerisation is such that the deposited layer remains a liquid long enough to allow it to flow. As the deposition proceeds and the film grows thicker, surface tension causes the layer to self planarize. The process thus achieves both gap-fill and planarization simultaneously.

Following deposition the layer is subjected to an in-situ heat treatment step in order to promote further polymerisation reactions and convert the material from the liquid form into a stable amorphous structure. An ex-situ furnace anneal (400°C , 30 minutes, N_2 ambient) is then carried out in order to effectively remove any residual moisture that remains as a by product of the polymerisation reactions.

The deposition of the low-k Flowfill[®] films was carried out using a Planar 200 dielectric deposition tool [1].

PHYSICAL PROPERTIES

Fourier transform infrared (FTIR) analysis revealed the presence of both an Si-C peak at 1275 cm^{-1} and a C-H peak at 2970 cm^{-1} for the methyl doped Flowfill[®] layers as shown in Figure 3. This confirms the incorporation of the CH_3 group bound to the silicon atom within the silica network.

The thermal stability for the material has been evaluated using FTIR and thermal desorption spectroscopy (TDS) measurements. Figure 5 shows the SiC/SiO and CH/SiO integrated peak area ratios, measured by FTIR, vs. annealing temperature. It can be seen that both the Si-C and C-H bonds remain stable up to annealing temperatures of around 500°C . Figure 6 shows typical thermal desorption spectroscopy (TDS) data for a low-k Flowfill[®] layer. The data shows that desorption of CH_4 ($M/z = 16$) and CH_3 ($M/z = 15$) only occurs at temperatures greater than 500°C , with negligible desorption occurring below this temperature. The thermal stability exhibited by the low-k Flowfill[®] layer is therefore compatible with existing metallisation schemes.

INTEGRATION

In order to integrate the low-k Flowfill® layer into an inter-metal dielectric scheme it is necessary to deposit a base layer (or liner) prior to depositing the low-k Flowfill® layer. The base layer is a plasma deposited oxide layer which has been optimised to act as a moisture barrier, preventing the possibility of H₂O or H₂O₂ from the Flowfill® layer interacting with the underlying structure. The base also provides a consistent adhesion layer to the overlying low-k Flowfill® layer. An oxide capping layer is then deposited on top of the low-k Flowfill® in order to provide mechanical stability during the final anneal step.

This process sequence gives rise to several integration issues:

Base layer dependence.

The use of a separate liner and capping layer will clearly contribute to the overall dielectric constant of the integrated three layer stack, since the individual layers will act as capacitors in series. The capacitance for an individual layer is given by:

$$C = \frac{k\epsilon_0 A}{t} \dots\dots\dots$$

(3)

where k = dielectric constant, ϵ_0 = permittivity of free space,
 A = capacitor plate area, t = thickness of dielectric.

and the overall capacitance of the integrated stack is given by:

$$\frac{1}{C_{total}} = \frac{1}{C_{liner}} + \frac{1}{C_{lowkFlowfill}} + \frac{1}{C_{cap}} \dots\dots\dots$$

(4)

It has been found by experiment that the total capacitance of the three layer stack incorporating the low-k Flowfill® is greater than that given by equation (4). Further investigations have shown that the dielectric constant of the low-k Flowfill® layer is actually dependant on the type of liner it is deposited onto. Low-k Flowfill® layers deposited onto silicon substrates exhibit a dielectric constant of 2.9, however the same film deposited onto a base layer exhibits a dielectric constant of ~3.4. A difference was also observed for low-k Flowfill® films deposited onto a nitride, a silane based oxide and TEOS based oxide which gave a dielectric constant of 3.7, 3.1 and 3.1 respectively. FTIR analysis has shown that the carbon content of the layers is independent of the liner used. This suggests that either the physical structure of the low-k Flowfill® layer has been affected, or that an interfacial effect has been introduced. As the Flowfill® deposition mechanism involves a surface reaction it is reasonable to assume that the condition at the surface of the liner may give rise to either effect. The exact mechanism is currently under investigation.

Since the base layer is required as a moisture barrier, a proprietary surface treatment step has been developed to minimise this effect. The use of the treatment step prior to the low k

Flowfill® deposition reduces the dependence of dielectric constant on the liner and hence the capacitance of the three layer stack to that predicted by equation 4 (Figure 7).

Degassing properties.

As described in a previous section, the low-k Flowfill® layer is subjected to a heat treatment step following deposition using a similar process to that used for the standard $\text{SiH}_4/\text{H}_2\text{O}_2$ Flowfill® process. This promotes polymerisation of silanols to an amorphous material. TDS has shown that this heat treatment step is insufficient for the low-k Flowfill® layers resulting in incomplete polymerisation. This can give rise to degassing of moisture during subsequent processing steps. The heat treatment step has therefore been optimised resulting in degassing properties similar to those of typical plasma deposited TEOS layers (Figure 8).

The resolution of these issues has allowed the low-k Flowfill® layer to be successfully integrated into 0.25µm technologies [7].

DIELECTRIC CONSTANT OPTIMISATION

The reduction in dielectric constant to 2.9 for the low-k Flowfill® layer has been attributed to the incorporation of the CH_3 group bound to the silicon atom in the oxide lattice. To determine the relationship between dielectric constant and the CH_3 content of the layers the carbon content has been determined using elastic recoil detection analysis (ERDA) and rutherford backscattering spectroscopy (RBS). The dielectric constant was determined by fabricating metal-oxide-metal capacitor structures and measuring their capacitance ($f = 1\text{MHz}$). Figure 9 shows the relationship between dielectric constant and carbon content for the low-k Flowfill® layers. The data was generated by varying the $\text{SiH}_4:\text{CH}_3\text{-SiH}_3$ gas flow ratio from 1:0 to 0:1. It can be seen that the dielectric constant is dependant on the amount of carbon and hence the number of CH_3 groups present in the layer. It can also be seen that the maximum carbon content that can be achieved repeatably is only ~ 12 at.% which is less than that predicted by the proposed reaction mechanism.

A possible explanation for the lower carbon content is that some of the Si-CH_3 bonds may have been broken as a result of the surface reaction between the $\text{CH}_3\text{-SiH}_3$ and H_2O_2 . It is possible to achieve a higher carbon content and hence a lower dielectric constant by varying the process conditions, as previously suggested [6], however these process conditions currently give rise to non-repeatability issues. The carbon content can be increased by increasing the process pressure or by decreasing the substrate temperature. The non-repeatability issues arise because both of these changes move the process towards the condensation point of the H_2O_2 precursor resulting in process instability. Current work is centred on further optimising the process conditions to allow a higher carbon content to be achieved.

NEW CHEMISTRIES

An alternative approach to lower the dielectric constant of the low-k Flowfill[®] films is to modify the process chemistry rather than the process conditions in order to include more carbon. Feasibility studies have shown that this can be achieved by using additional or alternative pre-cursor materials, some of which will now be considered.

Di-methyl silane $(\text{CH}_3)_2\text{-SiH}_2$ is a gaseous pre-cursor that contains two CH_3 groups bound to the silicon atom as opposed to one CH_3 for methyl silane. Di-methyl silane has been used in addition to methyl-silane to produce a layer with a dielectric constant of ~ 2.75 and properties substantially unchanged from those of the standard methyl-silane process. Matsuura et. al. [7] has taken this process further and shown that it is applicable to sub $0.25\mu\text{m}$ devices using both W plug and Forcefill[®] [8] Al plug processes. Device reliability was demonstrated using a $0.25\mu\text{m}$ two level interconnect scheme. Low via resistance, leakage current and Kelvin resistance, were achieved.

The methyl silane and di-methyl silane based films both result in carbon being bound into the oxide lattice via a Si-CH_3 bond which results in the termination of the siloxane chain. The addition of further CH_3 groups bound to the silicon atom is thought to be prohibitive because an increase in the number of CH_3 groups reduces the number of sites available to form the siloxane chain.

It is possible to incorporate carbon as part of the siloxane chain itself so that the siloxane chain is not terminated. Methylenebis-silane $(\text{SiH}_3\text{-CH}_2\text{-SiH}_3)$ is a gaseous pre-cursor that contains a CH_2 group bound between two SiH_3 groups. The SiH_3 functional groups can take part in the polymerisation reactions with H_2O_2 leaving the $\text{Si-CH}_2\text{-Si}$ backbone intact as part of the siloxane chain. This precursor has been used in place of methyl-silane to form a layer with a dielectric constant of ~ 2.7 .

An alternative approach is to utilise groups other than $-\text{CH}_3$ bound to the reactive $-\text{SiH}_3$ group. By correctly choosing the precursor material dielectric constants in the range 2.25-2.75 are achievable.

The use of alternative precursor materials has demonstrated that the low-k Flowfill[®] technology can be extended to achieve lower dielectric constants. These results are summarised in Figure 10.

SUMMARY

Optimisation of the $\text{CH}_3\text{-SiH}_3/\text{H}_2\text{O}_2$ low-k Flowfill[®] process has produced a stable low-k material with a dielectric constant of 2.9, and physical properties that should allow it to be integrated into current metallisation schemes.

Furthermore it has been shown that this CVD technology has the potential to produce a range of new oxide based materials with dielectric constants ≤ 2.5 by utilising various organic precursors.

ACKNOWLEDGEMENTS

The authors would like thank K.Giles and the CVD engineering team at Trikon Technologies for their invaluable contributions to this work, and also D.Moore and Dr. P.Timms at the University of Bristol, U.K., for their helpful discussions.

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- [2] M.Matsuura et al., IEDM, Dec. 11-14th 1994, pp117-120.
- [3] M.Matsuura et al, "An advanced planarizing interlayer dielectric using SiH₄ and H₂O₂ chemistry", 1995 Symposium on Dry Process.
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- [5] F.Gaillard et al, "Hydrogen peroxide and silane CVD process for pre-metal device applications", DUMIC, Feb. 20-21 1996
- [6] S.McClatchie et.al. "Low Dielectric Constant Flowfill[®] Technology for IMD Applications", DUMIC, Feb. 10-11th 1997.
- [7] M.Matsuura et. al., A Reliable Self-planarizing Low-k Intermetal Dielectric for Sub-quarter Micron Interconnects", IEDM, Dec. 7-10th, 1997.
- [8] A.J.McGeown, "High Pressure Aluminium Fill", ULSI 95, pp. 635-643.



Exhibit B

ULTRA-LOW DIELECTRIC CONSTANTS (ULDC) MATERIALS FOR MICROELECTRONICS AND ETEXTILES

UCLA Technology Available For Licensing

UCLA researchers in the Department of Mechanical and Aeronautical Engineering have developed and reduced to practice a low dielectric material composed of polymers infused with nanoparticles. The material can be used as an interlayer dielectric for large scale integrated interconnects. The material is very thin and has potential application for large area sensing and actuating systems utilizing woven electronic circuits.

Background:

Intensive research is being conducted on low dielectric constant materials. These include fluorinated silica glass, air gap formation, in fluorinated polymers, polyarythers, inorganic-organic hybrids, porous polymers and other materials. These exhibit dielectric constants in the range of 1.8 to 2.9.

Innovation:

Using specific formulation of polymers and nanoparticles infused into the polymer, a very thin material, on the order of 100 microns in thickness can be fabricated which produces a dielectric constant less than 1.1 at 100 KHz for FE-8% and GNP 4%. Other material systems have been tested and show very low dielectric constants versus competing materials.

The invention makes use of nanofiber technology to create an intense 3-D fibrous network resulting in a fully covered membrane with enormous pore surfaces. The dielectric and mechanical properties of the material can be tailored by selective formulation of the nanoparticle and polymer. The resultant material is produced using electrospinning.

Development to Date:

The dielectric material can be produced in sample quantities and are available for commercial testing. Many combinations of material systems have been processed and tested with the following results all at 100 KHz:

| <u>Material</u> | <u>Dielectric Constant</u> |
|------------------------|-----------------------------------|
| FE 8% | 1.08 |
| GNP 4% -A | 1.07 |
| GNP 4% -B | 1.14 |
| GNP 2% -B | 1.16 |
| PAN FE 4% | 1.14 |
| FE 1% | 1.16 |
| PAN FE 6% | 1.16 |
| GNP 2% A | 1.18 |
| PAN FE 2% | 1.28 |
| PAN (A) | 1.46 |
| PAN (B) | 1.24 |

Reference: UCLA Case No. 2003-039

**For information on licensing this invention,
please contact the office below.**

Office of Intellectual Property Administration
University of California, Los Angeles
10920 Wilshire Blvd., Suite 1200
Los Angeles, CA 90024-1406
Tel: 310-794-0558 Fax: 310-794-0638
email: ncd@resadmin.ucla.edu

NCD URL: <http://www.research.ucla.edu/tech/ucla03-039.htm>

Lead Inventor: Thomas Hahn

Search! Research Interests

UCLA Technologies Available for Licensing
<http://www.research.ucla.edu/tech>

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keywords: therapeutics prophylactic uclancd ucla technologies intellectual property patents technology transfer invention business card

Flowfill® Technology

Low-Dielectric-Constant Materials

WEI-JEN HSIA, WILBUR CATABAY, DUNG-CHING PERNG, & PETER J. WRIGHT, LSI Logic, Santa Clara, CA, USA
LIAM CUNNANE, KNUT BECKMANN, SIMON MCCLATCHIE & ADRIAN KIERMASZ, Trikon Technologies Ltd, Newport, Gwent, UK

ABSTRACT

The approach to the integration of low- k materials studied in this article is to use an inorganic material such as silicon dioxide doped with organic components. Embedded and non-embedded integration schemes are described. Electrical data shows that the low- k material performs as well as or better than a standard oxide.

INTRODUCTION

The integration of organic low-dielectric-constant materials presents the industry with many difficulties. Problems vary depending on the low- k material in question, but can include poor adhesion, issues with etching and resist removal creating the potential for via poisoning, and also thermal-conductivity problems with the more esoteric low- k films.

The approach studied in this article is to use an inorganic material such as silicon dioxide doped with organic components. The organic content reduces the dielectric constant but the inorganic backbone maintains film properties similar to that of standard silicon dioxide to minimize any integration problems.

CHEMISTRY

The low- k flow layer is formed through a condensation reaction between hydrogen peroxide and methylsilane [1], and the addition of silane can be used to control the k value. In this way the amount of carbon incorporated into the predominantly inorganic silica film can be adjusted. The maximum carbon content is achieved by complete removal of the silane from the reaction. Water is formed as part of the condensation reaction and a final anneal between 400 and 450°C ensures outgassing and the formation of a high-quality methyl-doped silicon dioxide film.

The low- k film exhibits good thermal stability, where carbon loss does not occur until ~ 500°C. Good adhesion is routinely achieved to a variety of materials including Ti, TiN, SiO₂, SiN, and Si. Low stress ~ $<5 \times 10^8$ dynes/cm² and unprecedented gap fill properties are also key attributes. Direct chemical-mechanical polishing (CMP) has been carried out with excellent results [2, 3].

Dielectric constant values as low as 2.7 can be achieved with standard methyl-based chemistry and are directly dependent on the carbon content in the film [1].

INTEGRATION SCHEMES

The Trikon advanced planarization layer typically consists of a plasma-deposited base or liner followed by a low-temperature flow layer, a plasma-deposited cap, and finally a furnace anneal (B-FF-C-FA). The exact functions of the base and cap layers have been reported elsewhere [4]. It is also possible to use a process flow where the Flowfill® is deposited thicker and the cap layer is replaced by a simple treatment step (B-FF-T), hence allowing a lower k value to be realized plate to plate as well as line to line; although this is still being investigated to fully understand the process advantages, there are obvious throughput and cost-of-ownership gains. The treatment step, its associated advantages, and film quality aspects will be discussed in a future article. There is some evidence that the base liner layer can be deposited ex situ or indeed possibly omitted, depending on the aluminium passivation and resist strip used during the metal etch (FF-T).

Taking these basic process steps into account, a number of possible integration schemes have been considered, as follows.

EMBEDDED

The embedded approach essentially uses a B-FF-C-FA or B-FF-T sequence deposited over the metal layer. Using either direct CMP or a sacrificial oxide, the structure is CMP'd back to the metal and then capped with a standard oxide. The vias are then etched through the standard oxide layer using usual oxide etch chemistries and all issues to do with low- k via poisoning are eliminated. This is clearly not a new idea and has been used with spin-on materials for some time. It does, however, allow a low- k CVD film to be used between metal lines without the added difficulty of low- k via integration.

NON-EMBEDDED

Both B-FF-C-FA and B-FF-T sequences can be used with this scheme. These process schemes are deposited directly onto the metal structure and vias are etched through the complete stack. Depending on the device metallization, it is possible to use direct or sacrificial polishing exactly as with the embedded approach. Owing to the planarization capability of Flowfill®, it is also possible to delay the use of polishing, or reduce the number of polishing steps in a device. Data has been presented elsewhere which shows improved uniformity, higher throughputs, and reduced CMP consumables when polishing Flowfill, because of its planar nature and the simple fact that less polishing is required to produce the final flat surface [3].

Using a non-embedded approach requires specific expertise and know-how on how to prevent via poisoning in low- k films. This can be an expertise in the strip and liner

Figure 1
Scanning electron micrograph
of the integrated metal scheme

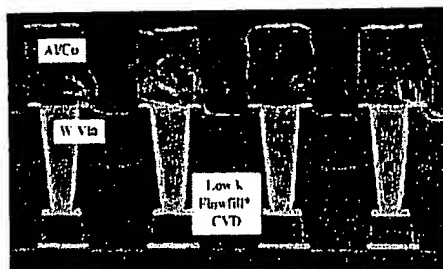
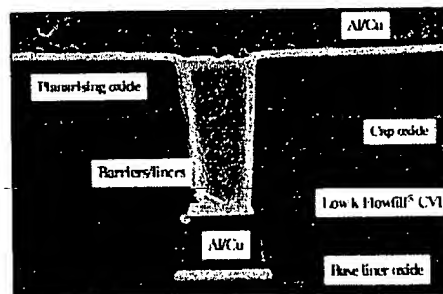


Figure 2
Typical cross-section through a via



technology or adopting a more complex integration scheme where a hard mask is used to pattern the via, thereby avoiding the use of resist strip chemistries. Typically, the lower the dielectric constant of the low-*k* material, the more difficult it is to prevent via poisoning unless a more complex integration scheme is used. A key advantage of a non-embedded approach is that the low-*k* material can be realized both between the lines and between the metal levels. Clearly a B-FF-T sequence with direct CMP will offer more advantages with respect to *k* value and cost-of-ownership issues than a B-FF-C-FA sequence with a sacrificial CMP layer.

With either the embedded or the non-embedded approach, the base liner can be used ex situ or eliminated as explained above, offering a direct-on-metal (DOM) process.

Figure 1 shows a scanning electron micrograph of the integrated metal scheme. Throughout the work, tungsten plug was used for the via and aluminum/copper alloy was used for the interconnect; 0.18 μm design rules were used, although exact dimensions are proprietary at this stage. Various aspect ratios of both the vias and the metallization in the device can be determined to indicate the complexity of the integration.

In general terms the IMD (Inter Metal Dielectric) process flow was as follows:

1. deposit low-*k* IMD sequence
2. CMP
3. lithography
4. via etch/resist strip/clean
5. degas/deposit liners/W plug
6. W CMP
7. deposit Al
8. lithography
9. etch/resist strip/clean.

Although at first sight the actual dielectric portion of the scheme looks complex, it is in fact relatively straightforward since a B-FF-C or B-FF-T sequence is carried out completely in situ using a predefined process recipe. This is particularly important for ASIC devices, where fast turn-around and minimum process delay times are important. From a

processing point of view, only one vacuum break occurs during the complete dielectric deposition; this is between the cap oxide and the planarizing sacrificial oxide if one is used.

Figure 2 shows a typical cross-section through a via. The base-low-*k* Flowfill[®]-cap structure can be clearly observed. Low-*k* material can clearly be seen both between the metal lines and part way up the via, highlighting the fact that this process is a true non-embedded approach.

ELECTRICAL DATA

Wherever possible, electrical data for low-*k* Flowfill[®] has been compared to a standard oxide, produced by a high-density plasma (HDP) source. In all tests the low-*k* material performed as well as or better than the standard. The singular property which was always better was the capacitance of the IMD layer.

Apart from its gap fill capability, the *k* value of low-*k* Flowfill[®] can be varied according to the requirements of the device and its design criteria. This offers the possibility for existing devices that use a standard oxide to operate faster, provided the design criteria allow the speed to be realized. The value of dielectric constant can be easily varied from that of HDP oxide ($k \sim 4.2$) to sub-3.0 values. This is done simply by changing the process conditions; no extra hardware is required.

Figure 3 shows the performance of a Kelvin via compared to an HDP oxide.

Figure 4 shows a via chain plot for 0.25 μm vias, again comparing low-*k* Flowfill[®] to HDP. There is no via poisoning evident, nor indeed any evidence of any high-resistance contacts.

CONCLUSIONS

A flexible low-*k* CVD film has been integrated into a device with 0.18 μm design rules. The physical and electrical characteristics are excellent.

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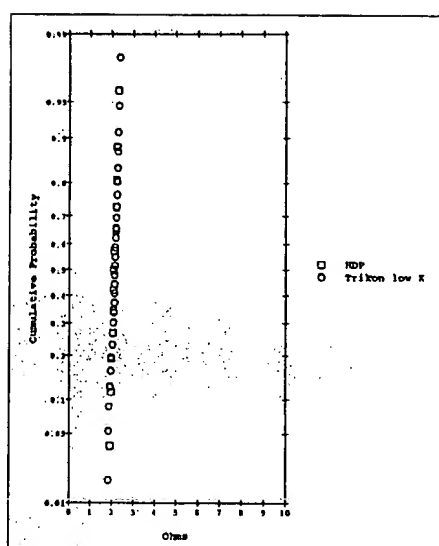
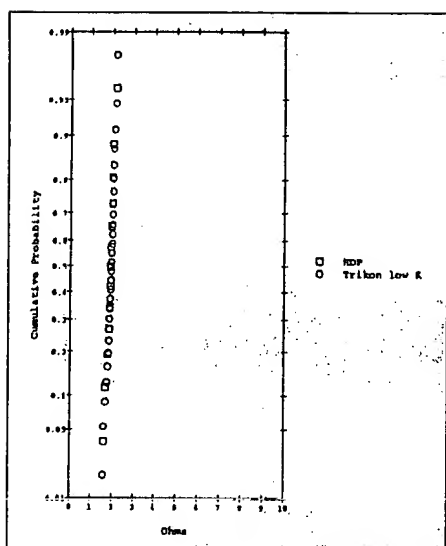


Figure 3 (far left)
Electrical data for a Kelvin via
compared to an HDP oxide

Figure 4 (left)
Via chain plot for 0.25 μ m vias

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[Reader Ref. 38]

Origin of low dielectric constant of carbon-incorporated silicon oxide film deposited by plasma enhanced chemical vapor deposition

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Dielectric properties of carbon-incorporated silicon oxide (SiOC) films deposited by plasma enhanced chemical vapor deposition (PECVD), using a bis-trimethylsilylmethane precursor, were compared with the dielectric properties of silicon oxide thermally grown in a furnace (thermal oxide) and also with oxide deposited by PECVD using a tetrathoxysilane (TEOS) precursor (PE-TEOS oxide). The electronic contribution to the dielectric constant of the three oxide films was calculated from their refractive indices measured at 632.8 nm by ellipsometry. Ionic contributions were computed from their IR reflection spectra, measured at 650–4000 cm^{-1} , by using the Kramers–Kronig relation. The dipolar contribution was qualitatively analyzed from temperature dependence of the polarizability, on a per unit volume basis. The dielectric constant of the SiOC films, which was measured at 1 MHz, decreased from 4.2 to 2.3 as the carbon content increased from 0 to 19.6 at. %. Although there was a significant reduction of the dielectric constant, the electronic contribution was only slightly changed from 2.10 to 1.90, whereas the ionic polarization noticeably changed from 1.86 to 0.25. SiOC films showed considerable dipolar contribution when compared to thermal oxide and PE-TEOS oxide films, but the dipolar contribution in the SiOC films became negligible as the carbon content was increased. The variation of each contribution with the carbon content shows that carbon incorporation leads to a decrease in the electronic and ionic contributions. The reduction of the ionic contribution was the predominant factor leading to a decrease in the overall dielectric constant. © 2001 American Institute of Physics.

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I. INTRODUCTION

As the design rule of integrated circuit (IC) devices approaches 0.1 μm and beyond, the interconnection delay (RC delay) begins to govern overall device delays. In addition, the shrinkage of device dimensions causes cross talk between signal paths and higher power consumption. To realize high performance in IC devices it is necessary to reduce the RC delay, cross talk, and power consumption. An effective method to make these reductions is to decrease the parasitic capacitance by employing a low dielectric constant (low- k) material in the intermetal dielectrics (IMD).^{1,2} Over the past few years, fluorinated SiO_2 (SiOF)³ films and hydrogen silsesquioxane (HSQ) have been studied for this application, but it has been reported that they are limited to reducing the dielectric constant below 3.0.^{2,4}

Recently, various organic polymers and carbon-based composite materials were extensively investigated for low- k IMD materials as an alternative to conventional SiO_2 . Among the materials investigated, carbon-incorporated silicon oxide (SiOC) is considered a promising candidate in the 0.13 μm technology and beyond.⁵ Previously we reported

that it is possible to reduce the dielectric constant of a SiOC film to 2.4 and that the film is thermally stable up to 500 $^{\circ}\text{C}$.⁶ Korczynski⁷ and Pang⁸ reported that the low dielectric constant of a SiOC film resulted from hydrocarbon (methyl group, $-\text{CH}_3$) incorporation and the film's low density. However, the exact mechanism to reduce the dielectric constant of SiOC films has not been clearly understood until now.

In this study the origin of the low dielectric constant of SiOC films was investigated by examining the dielectric contribution of electronic, ionic, and dipolar polarization. Each contribution of these polarization modes was also investigated in thermally grown silicon dioxide (thermal oxide) and silicon oxide deposited by plasma enhanced chemical vapor deposition (PECVD) while using a tetrathoxysilane precursor (PE-TEOS oxide) for comparison.

II. EXPERIMENT

A. Thin film preparation and characterization

SiOC films were deposited on p -type (100) Si substrates by PECVD using a bis-trimethylsilylmethane (BTMSM) liquid precursor. The PECVD system used in this study and the detailed deposition conditions have been described elsewhere.⁶ Thermal oxides were grown by dry oxidation of

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the Si substrate at 1 atm and O₂/N₂ ambient conditions. The tetrathoxysilane precursor (PE-TEOS) oxide films were deposited by PECVD using a tetraethoxysilane (TEOS) precursor. The film thickness and refractive index were measured by ellipsometry, at a wavelength of 632.8 nm (model: Gaertner L116-HP85B) and spectroscopic ellipsometry (model: Sopra ESG). The chemical bonding state and the dielectric constant in the IR region of the SiOC film were characterized by using Fourier transform infrared (FTIR) spectroscopy. This was operated in the absorbance and reflectance mode with a Bio-Rad FTS40. The film composition was determined by elastic recoil detection time-of-flight (ERD-TOF) analysis. The film density was directly calculated from the ratio of mass to volume of the film. The film mass was measured by an electronic balance (model: AND HM202) with a resolution of 0.01 mg. The film volume was calculated by three-dimensional mapping of the film thickness, measured at 625 different points on the 4-in. wafer. The dielectric constant at 1 MHz was obtained by a HP4280A C-V meter on the metal-insulator-semiconductor (MIS) structure.

B. Analysis of dielectric polarizations

The dielectric constant is a frequency-dependent, intrinsic material property. The constant can be divided into three components that result from electronic, ionic, and dipolar polarization. The dielectric constants measured at 1 MHz consist of electronic ($\Delta\epsilon_e$), ionic ($\Delta\epsilon_i$), and dipolar ($\Delta\epsilon_d$) contributions, as expressed in

$$\epsilon_r(\text{at } 1 \text{ MHz}) = 1 + \Delta\epsilon_e + \Delta\epsilon_i + \Delta\epsilon_d. \quad (1)$$

The dielectric constant of a material can also be calculated from the refractive index, as expressed in

$$\epsilon_r(\lambda) = n^2(\lambda) + k^2(\lambda), \quad (2)$$

where ϵ_r is a relative dielectric constant, n is a real part of a refractive index, k is an imaginary part of a refractive index, and λ is the wavelength of a light source.

The pure electronic contribution to the dielectric constant ($\Delta\epsilon_e$) was calculated from the refractive index obtained in the visible-ultraviolet (vis-UV) region. Since the extinction coefficient k of SiO₂-based materials is normally negligible in this region, the relative dielectric constant in Eq. (2) can be simply expressed as n^2 .^{9,10}

The ionic contribution to the dielectric constant ($\Delta\epsilon_i$) was calculated by subtracting the dielectric constant at 632.8 nm, $1 + \Delta\epsilon_e$, from the dielectric constant in the IR region, $1 + \Delta\epsilon_i + \Delta\epsilon_d$. To obtain the dielectric constant in the IR region, the refractive index was calculated from the Kramers-Kronig dispersion relation.

The original Kramers-Kronig relation is expressed as in¹⁰⁻¹³

$$n_i = 1 + \frac{\pi}{2} P \int_0^\infty \frac{\nu k(\nu)}{\nu^2 - \nu_i^2} d\nu, \quad (3)$$

where n_i is the real part of the refractive index at the i th wave number, P is the principal value ensuring integral from zero to infinity, ν_i is a wave number at i , and k is the imaginary part of the refractive index (extinction coefficient).

The integration in Eq. (3) cannot be performed from zero to infinity since the IR spectrum is obtained only in a finite region (650–4000 cm⁻¹). By introducing the refractive index at 632.8 nm (n_e , $\nu = 15\,800$ cm⁻¹), the above integration is approximated in a finite region of the IR spectrum as is shown in Eq. (4). The dielectric constant in the range from 4000 cm⁻¹ to infinity is approximated as the dielectric constant at 632.8 nm calculated from the refractive index by ellipsometry, as explained in Eq. (2).

$$n_i \approx n_e + \frac{\pi}{2} \left[\int_{650}^{\nu_i} \frac{\nu k(\nu)}{\nu^2 - \nu_i^2} d\nu - \int_{\nu_i}^{4000} \frac{\nu k(\nu)}{\nu^2 - 15\,800^2} d\nu \right], \quad (4)$$

where n_e is the refractive index at 632.8 nm (15 800 cm⁻¹).

The IR reflectance spectra of the films were obtained from FTIR spectroscopy (Bio-Rad FTS40). The commercial program WIN-IR™ (from Bio-Rad) was used to transform these reflectance spectra into IR region refractive indices. The IR region dielectric constants were calculated by Eqs. (2) and (4). However, the ionic contribution obtained from this transformation may not be quantitatively precise due to some approximations used in the transformation calculation and the restricted instrumental resolution of FTIR spectroscopy. To manage this lack of precision the IR region dielectric constants were corrected by normalizing with the thermal oxide ionic contribution ($\Delta\epsilon_i$). The thermal oxide value $\Delta\epsilon_i$ was obtained from Eq. (1), where $\Delta\epsilon_d$ is zero. ($1 + \Delta\epsilon_e$) is the same as the square of the refractive index at 632.8 nm, and ϵ_r is obtained from the C-V measurement at 1 MHz.

The dipolar polarization was qualitatively estimated by the Clausius-Mosotti relation in^{9,14}

$$\left[\frac{(\epsilon_r - 1)}{(\epsilon_r + 2)} \right] = \frac{\rho}{M} \times \frac{N_A}{3\epsilon_0} (\alpha_{ie} + \alpha_d), \quad (5)$$

$$\alpha_d = \frac{p^2}{3kT} (\text{dipolar polarizability}),$$

where M is molecular weight, ρ is density, ϵ_r is the dielectric constant at 1 MHz, ϵ_0 is the vacuum dielectric constant, α_{ie} is the electronic and ionic polarizability, α_d is the dipolar polarizability, p is the dipole moment, k is the Boltzmann constant, and T is absolute temperature.

Unlike electronic and ionic polarizabilities, dipolar polarizability is inversely proportional to temperature, as is shown by Eq. (5). Therefore the temperature dependence of the polarizability per unit volume can imply that a dipole moment exists in the film. In this study, this dependence was investigated by varying the measurement temperature from 20 to 180 °C.

III. RESULTS AND DISCUSSION

Thermal oxide film was grown in a dry oxidation atmosphere at 1000 °C. PE-TEOS oxide and SiOC films were deposited on *p*-type Si(100) by PECVD. The carbon content incorporated into the films was controlled by the substrate temperature. The basic physical properties of the films are summarized in Table I.

TABLE I. Physical properties of thermal oxide, PE-TEOS oxide, and SiOC films having varying carbon content.

| | Thermal oxide | PE-TEOS oxide | SiOC | | |
|-------------------------------------|---------------|---------------|---|---|---|
| | | | low | mid | high |
| film thickness (nm) | 105.4 | 103.8 | 100.0 | 100.0 | 99.8 |
| refractive index (at 632.8 nm) | 1.458 | 1.462 | 1.435 | 1.396 | 1.360 |
| film density (g/cm ³) | 2.40 | 2.24 | 2.09 | 1.59 | 1.17 |
| porosity (%) ^a | 0 | NA | 13.6 | 23.6 | 39.0 |
| carbon content (at. %) ^b | NA | NA | 5.5 | 10.5 | 19.6 |
| [chemical composition] | | | [SiO _{1.9} C _{0.17}] | [SiO _{1.3} C _{0.27}] | [SiO _{0.9} C _{0.46}] |

^aPorosity (%) was calculated from the film density and molar weight of each composition, based on the assumption that the thermal oxide has no porosity.

^bat. % (from ERD-TOF)=100×[C]/([Si]+[O]+[C]).

Figure 1 shows the variation of film densities and refractive indices measured at 632.8 nm as a function of film carbon content. The refractive index of the PE-TEOS oxide film was slightly higher than that of the thermal oxide film. This difference is due to the two films' composition differences. As more carbon was incorporated into the SiOC film, both the film density and refractive index decreased. Previously, we reported that the decrease of film density in SiOC films resulted from termination of the O–Si–O bonding network by replacing oxygen atoms with hydrocarbon groups (–CH_n).⁶ The SiOC film density was reduced to half the value of the PE-TEOS oxide density when the carbon content was increased to 19.6 at. %, while the refractive index slowly decreased from 1.46 to 1.36. For “aerogel” and “xerogel,” which are low dielectric constant materials having the same composition as PE-TEOS oxide of SiO_{2-x}, their refractive indices are about 1.20 when their film densities are half that of the PE-TEOS oxide.¹⁵ Considering that the incorporation of hydrocarbon groups (–CH_n) altered the SiOC film stoichiometry, as shown in Table I, the relatively small refractive index decrease was attributed to the change of SiOC film's molar electronic polarizability. This change resulted from the compositional change.

The dielectric constants measured at 1 MHz and the electronic dielectric constants obtained from the squares of the refractive indices at 632.8 nm are shown in Fig. 2. The

dielectric constants at 1 MHz declined drastically with increased carbon content of the SiOC film. The decrease of the electronic contribution ($\Delta\epsilon_e$) to the dielectric constant was quite small in comparison with the variation of the dielectric constant at 1 MHz. From these results we found that either ionic ($\Delta\epsilon_i$) or dipolar ($\Delta\epsilon_d$) contributions to the dielectric constant were the dominant factors to reduce the dielectric constant of the SiOC film, rather than that from the electronic contribution ($\Delta\epsilon_e$). However, there is no information on the separate values of each $\Delta\epsilon_i$ or $\Delta\epsilon_d$, as only their combined contribution to the dielectric constant measured at 1 MHz is available.

To examine the amount of ionic contribution to the SiOC film's dielectric constant we calculated the dielectric constants in the IR region. This was calculated from the IR reflectance spectra by using the Kramers–Kronig transformation method described in the previous section. Figure 3 represents the dispersive characteristics of the dielectric constants in the IR region of 650–4000 cm⁻¹. When the wavelength of the applied IR light exceeds about 1300 cm⁻¹, the ionic polarization can no longer follow the applied field and thus the dielectric constant is only composed of the electronic contribution ($1 + \Delta\epsilon_e$). In the region of 1000–1300 cm⁻¹, a resonant increase of the dielectric constant occurs

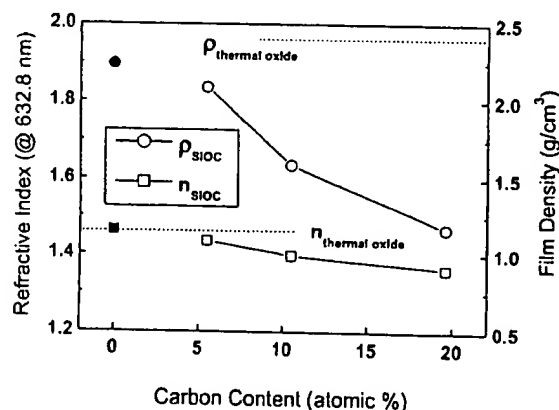


FIG. 1. Variations of film density and refractive index for the PE-TEOS film and the SiOC films as a function of carbon content.

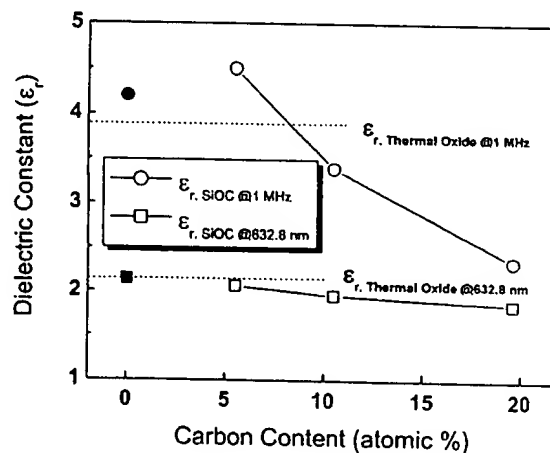


FIG. 2. Variations of dielectric constants, measured at 1 MHz, and electronic dielectric constants, calculated from the refractive index at 632.8 nm, for the PE-TEOS film, and the SiOC films as a function of carbon content.

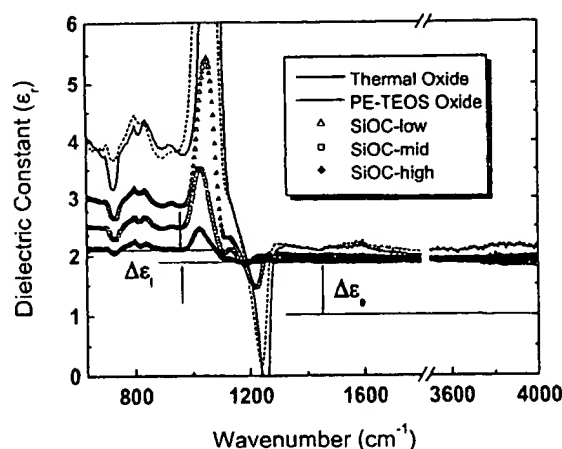


FIG. 3. Dielectric constants in the IR region calculated from the IR reflection spectra for the thermal oxide, PE-TEOS oxide, and SiOC films by using the Kramers–Kronig relationship. Magnitude of the dielectric constants was corrected by normalizing with the ionic contribution ($\Delta\epsilon_i$) of the thermal oxide (1.80).

due to IR absorbance by the Si–O stretching vibration mode (near 1100 cm^{-1}) in the SiOC film. The ionic contributions, due to other vibration modes in the SiOC film, are negligible because IR absorbances by other modes are considerably smaller than that by the Si–O stretching mode.^{10,16} At lower than the resonant frequency in the IR region, the dielectric constant contains the ionic contribution ($\Delta\epsilon_i$) as well as the electronic contribution.

As more carbon was incorporated in the SiOC films, the ionic contribution ($\Delta\epsilon_i$) to the dielectric constant decreased gradually, as depicted in Fig. 3. The reduced ionic contribution might be a consequence of a decrease of the Si–O bond in the film, which has strong ionic polarization.^{9,10} A previous study has shown that the Si–CH_n bond terminated the O–Si–O bonding network, causing a decrease in film density.⁶ We note that the ionic contribution's decrease can be explained by a decrease in film density. The replacement of Si–O by Si–CH_n bonds can also play a role in decreasing the ionic contribution due to the Si–CH_n bond being less ionic than the Si–O bond. With increasing degree of replacement, the oxygen deficiency in the film markedly increased, as shown in Table I. This led to a decrease of the ionic contribution due to the additional reduction of the Si–O

bond. Thus we can conclude that the decrease of Si–O bonds played a crucial role in reducing the measured ionic contribution to the dielectric constant. As is shown in Fig. 3, the lower intensity of the resonant peak (near 1100 cm^{-1}), for a film having higher carbon content, corresponds to a lower ionic contribution. This is good verification of our deductions.

Using Eq. (1), each contribution to the dielectric constant can be calculated from the dielectric constant measured at 1 MHz (ϵ_r), the refractive index (n), and the ionic contribution ($\Delta\epsilon_i$) of Fig. 3. Electronic, ionic, and dipolar contributions ($\Delta\epsilon_e$, $\Delta\epsilon_i$, and $\Delta\epsilon_d$) to the dielectric constant, with respect to the film carbon content, are summarized in Table II. The value of $\Delta\epsilon_d$ was obtained by subtraction of ($n^2 + \Delta\epsilon_i$) from ϵ_r . It should be noted that $\Delta\epsilon_i$ and $\Delta\epsilon_d$ are not exact values but are normalized values from $\Delta\epsilon_i$ (1.80) of the thermal oxide. This is based on the assumption that the thermal oxide has no dipolar polarization.

As mentioned above, the electronic contribution in the SiOC films was slightly decreased with increasing carbon content, but carbon content had little influence on reducing ϵ_r . On the other hand, the ionic contribution drastically decreased as more carbon was incorporated. This decrease in the ionic contribution is a predominant factor for the low dielectric constant of the SiOC films. However, the SiOC films showed considerable dipolar contributions when contrasted with the thermal oxide and PE-TEOS oxide. Since a [SiO₄] tetrahedron is symmetric, pure silicon dioxide does not have a dipolar polarization. When a hydrocarbon group (–CH_n) replaces an oxygen site, the [SiO₄] tetrahedron loses its symmetry by changing into [SiO₃(CH_n)]. The carbon incorporation, moreover, led to the oxygen-deficient SiOC film, as summarized in Table I. These configurational changes help explain the increased dipolar contribution in the SiOC films. Interestingly, even though carbon incorporation promotes dipole generation, the dipolar contribution of a film having a higher carbon content was lessened. This result can be explained by considering the film density. The SiOC film with a higher carbon content has a larger molar dipole moment than that with lower carbon content; nevertheless, its smaller density is considered to cause the reduction of apparent dipolar contribution.

Considering the Clausius–Mossotti relation [Eq. (5)], the dipolar contribution ($\Delta\epsilon_d$) could be qualitatively esti-

TABLE II. Dielectric constants in various frequency regions and contribution values for thermal oxide, PE-TEOS oxide, and the SiOC films having varying carbon content.

| | Thermal oxide | PE-TEOS oxide | SiOC | | |
|--|---------------|---------------|------|------|------|
| | | | low | mid | high |
| $1 + \Delta\epsilon_e^a$ | 2.13 | 2.14 | 1.97 | 1.96 | 1.90 |
| $1 + \Delta\epsilon_e + \Delta\epsilon_i^b$ | 3.90 | 3.97 | 2.96 | 2.52 | 2.15 |
| $1 + \Delta\epsilon_e + \Delta\epsilon_i + \Delta\epsilon_d^c$ | 3.90 | 4.20 | 4.50 | 3.39 | 2.34 |
| $\Delta\epsilon_e$ | 1.13 | 1.14 | 0.97 | 0.96 | 0.90 |
| $\Delta\epsilon_i$ | 1.77 | 1.83 | 0.99 | 0.56 | 0.25 |
| $\Delta\epsilon_d$ | 0 | 0.23 | 1.54 | 0.87 | 0.19 |

^a($1 + \Delta\epsilon_e$) is obtained from n^2 (at 632.8 nm).

^b($1 + \Delta\epsilon_e + \Delta\epsilon_i$) is obtained from Kramers–Kronig conversion of the IR reflection spectrum.

^c($1 + \Delta\epsilon_e + \Delta\epsilon_i + \Delta\epsilon_d$) is obtained from C–V measurement at 1 MHz.

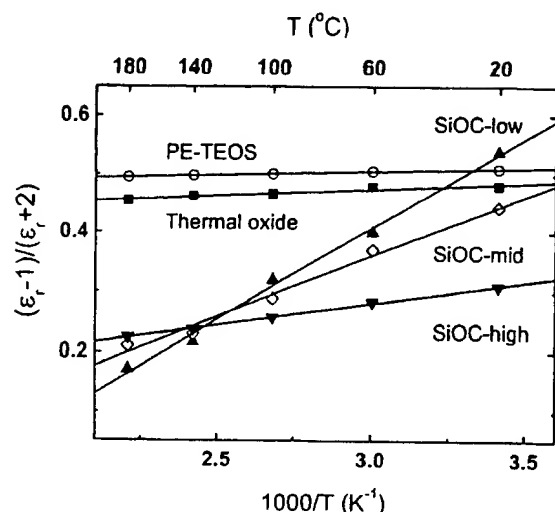


FIG. 4. Variations of the polarizability per unit volume for the thermal oxide, PE-TEOS oxide, and SiOC films as a function of measurement temperature.

mated from the temperature dependence of the polarizability per unit volume $[(\epsilon_r - 1)/(\epsilon_r + 2)]$. The variation of the dielectric constant (ϵ_r) due to measurement temperature and carbon content in the film was determined. Figure 4 shows the variation in the polarizability as a function of the temperature used in the $C-V$ measurements at 1 MHz. The measured polarizability of the SiOC film was inversely proportional to the temperature and its slope to $(1/T)$, which is proportional to the dipole moment per unit volume, and was gradually reduced with increasing carbon content of the film. Both the thermal oxide and PE-TEOS oxide films showed very little temperature dependence of their polarizability and from this the negligible dipolar contribution in each film^{10,16} could be confirmed. These results are in good agreement with the dipolar contribution values shown in Table II.

IV. CONCLUSION

The electronic, ionic, and dipolar contributions to the dielectric constants of the SiOC films were separately investigated using their frequency-dependent dielectric constants. The electronic contribution slightly decreased with increasing carbon content. This reduction of the electronic contribution can be explained by a decrease in the film density. How-

ever, the reduced electronic contribution had little influence on reducing the overall dielectric constant. The ionic contribution significantly decreased as more carbon was incorporated. This appears to be caused by decreased film density, replacement of Si-O by Si-CH₃, and increased oxygen deficiency of the film. On the contrary, the SiOC films showed considerable dipolar contributions as compared with the thermal oxide and PE-TEOS oxide. This dipolar contribution is considered to result from a configurational change of [SiO₄] tetrahedron due to carbon incorporation. However, the film having a high carbon content showed a negligible dipolar contribution as the decreased film density contributed to reducing the dipole concentration. Therefore the low dielectric constant of the SiOC films mainly results from reduction of the ionic contribution, which can be explained by the decrease of the Si-O bond in the film.

ACKNOWLEDGMENTS

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Building Copperopolis II

Using optical metrology to monitor low-k dielectric thin films

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Because controlling curing temperature and time is critical for producing high-quality spin-on dielectric thin films, rapid feedback from metrology tests on thermal process tools is needed to correct tool drift.

As device features of ULSI circuits continue to shrink, the capacitance of the interlevel dielectric (ILD) material becomes an increasingly limiting factor on the overall performance of these chips. An industrywide effort is under way to search for a next-generation low-k ILD material to replace traditional silicon dioxide. Many potential low-k materials belong either to the inorganic polymer family (e.g., organosilicates) or the organic polymer family.^{1,2} The molecular structure of the cross-link network in these polymers is critical because it affects their electrical, thermal, and mechanical properties as well as their optical properties, such as index of refraction (n) and extinction coefficient (k). Because n and k values depend both on the electronic structure of a polymer and on its porosity, they can serve as good indicators for monitoring low-polymer formation.

One low-k ILD material is SiLK from Dow Chemical (Midland, MI), a spin-on organic polymer that can be readily deposited using a conventional spin-coater. This dielectric thin film in its as-deposited state must undergo a thermal curing process to form a desired polymer structure and to achieve correct mechanical, thermal, and electrical properties. The curing process takes place at temperatures between 400° and 450°C in standard thermal-processing equipment (furnaces, ovens, or hot-plates). The cured resins have a dielectric constant of 2.65 and can withstand temperatures as high as 490°C. Their high thermal stability permits integration with current multilevel interconnect processes. However, because the polymerization of these resins is thermally activated, controlling the curing temperature and time is critical to producing high-quality cured films.^{3,4} Thus, it is essential to have immediate feedback from metrology tests on thermal process tools to detect and correct tool drift promptly. This article discusses optical metrology methods for monitoring cured low-k dielectric thin films.

Thin-Film Optical Metrology

One of the most important advantages of optical metrology arises from its nondestructive nature, which permits measurements on product wafers and active device areas. In-line optical metrology can monitor the performance of thermal processing equipment by conducting real-time measurements on product wafers, thus eliminating monitor wafers and minimizing rework or loss of product wafers as a result of out-of-control equipment.

Thin-film optical metrology provides fast and precise real-time measurements of thin-film thicknesses and optical constants. In semiconductor manufacturing, spectroscopic reflectometry and ellipsometry are wide

used optical metrology methods for measuring these parameters in-line. Thin-film metrology tools measure the reflectance or ellipsometric spectrum of thin films and extract values of thicknesses or optical constants.

Because there are a maximum of two independent optical measurement data (Ψ and Δ in spectroscopic ellipsometry) at each wavelength (λ), the maximum number of unknowns that can be determined in the whole spectrum is $2W$ (where W is the number of wavelengths). Materials with finite light absorption have two unknowns (n and k) at each wavelength and one additional unknown in the film thickness. Therefore, the total number of unknowns is double the number of wavelengths plus one—that is, $2W + 1$. Because the number of unknowns resulting from $2W + 1$ is at least one too many to be determined from available spectroscopic ellipsometry or spectroscopic reflectometry data, it is necessary to employ a dispersion model which describes the functional dependence of n 's and k 's on wavelength based on N fit parameters. Therefore, the total number of unknowns becomes $N + 1$. As long as $2W \geq N + 1$, film thickness and optical constants can be determined simultaneously by numerically iterating $N + 1$ variables to fit spectra.

The results of the research discussed in this article on the use of optical metrology to monitor a low- k resin curing process demonstrate that with a correct dispersion model, the $n(\lambda)$, $k(\lambda)$, and thickness values for a dielectric thin film can be determined from reflectance measurements. In the ultraviolet (UV) spectral region, the refractive indices values (RIs), or $n(\lambda)$ s, are found to correlate to the curing conditions. By measuring the variation of n , the degree of curing in the resin can be monitored. For example, at a wavelength of 314 nm, the RIs of dielectric thin films change systematically with the curing parameters of temperature and time.

Based on the relationship between the optical constants of resins and their curing parameters, a single-parameter empirical interpolation model for resin optical constants was developed. With this interpolation model, the curing of the resins can be readily monitored with an automated thin-film optical metrology tool that provides prompt feedback on the condition of the thermal processing equipment.

Sample Preparation and Instrumentation

For this study, a matrix of SiLK-I dielectric thin-film samples was prepared by varying the curing temperature and time. First, a spin-on process deposited resins on bare silicon wafers, which then went through a hard bake step at 310°C for 90 seconds and were subsequently cured on hot plates in a nitrogen ambient. The curing temperature varied from 400° to 470°C and the curing time from 30 to 360 seconds. The sample thicknesses were approximately 7400 Å.

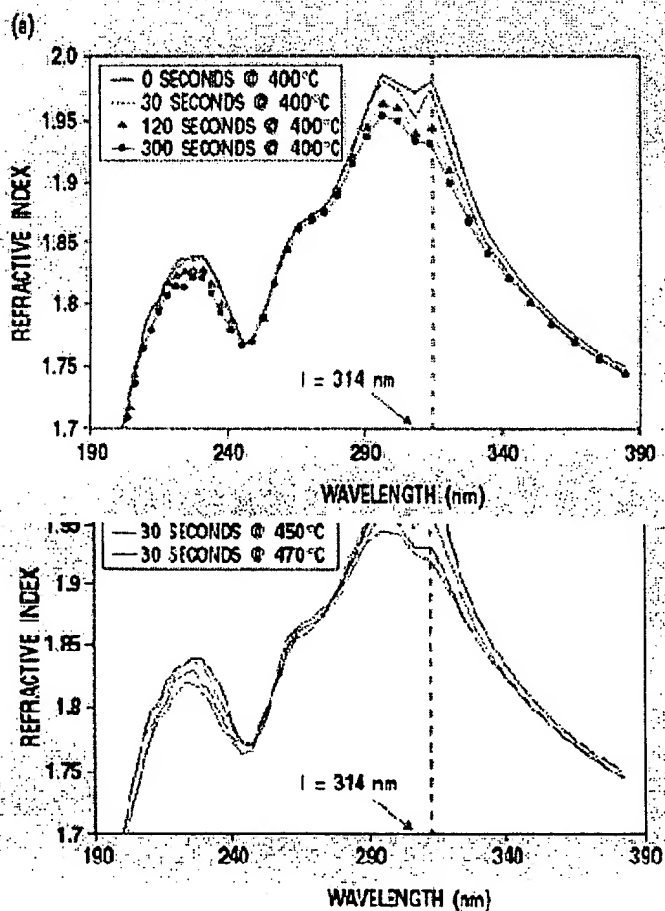
To determine the optical constants $n(\lambda)$ and $k(\lambda)$ of these samples, a variable-angle spectroscopic ellipsometer (VASE) from J. A. Woollam (Lincoln, NE) was used.⁵ A very powerful off-line analysis instrument for the characterization of the optical properties of thin films, this instrument utilizes a monochromator to control incident light wavelength so that only the light of a single wavelength is incident onto the sample during each measurement.⁶ It measures two ellipsometric data at each wavelength, Ψ and Δ , and completes the spectrum by scanning through all wavelengths. This way, the measured Ψ and Δ spectra truly represent the spectroscopic response of the sample. If a thin film's thickness is known, its $n(\lambda)$ and $k(\lambda)$ can be accurately determined by direct calculation from Ψ and Δ values.

In general, organic polymers are transparent or nearly transparent in the visible spectral region, from which their thicknesses can be extracted by fitting that part of a spectrum using a Cauchy dispersion formula. The

complicated optical constant response in the UV spectral region can then be determined directly from Ψ and Δ spectra.⁷ By choosing an appropriate dispersion model for the optical constants, the complicated n and k spectra from the VASE measurement results were parameterized. The production worthiness of the parametric dispersion model was tested by running it on a NanoSpec 8000XSE high-throughput thin-film metrology tool from Nanometrics (Sunnyvale, CA). This tool determines thickness values and the n 's and k 's of thin films by measuring and fitting spectroscopic reflectance spectra, spectroscopic ellipsometric spectra or a combination of both. Its modeling capability and data-fitting algorithm enable the analysis of data from a wide range of materials and layered structures while simultaneously determining thickness and optical constants.⁸

Single-Parameter Empirical Interpolation

Using the VASE measurements, a spectral window between 280 and 340 nm was found where the RIs of the thin-film samples changed monotonically with cure temperature and time. Figure 1 shows the measurements for UV spectra only. In the visible-wavelength region, RI variation was not significant. As demonstrated in both Figures 1a and 1b, sensitivity to the curing parameters of time and temperature was highest at a wavelength of 314 nm. The total magnitude of the RI change was 0.065 at this wavelength. Figure 2 plots RI for all test samples at 314 nm, denoted as $n(314 \text{ nm})$. Since the $n(314 \text{ nm})$ value decreases when either the curing time or temperature was increased, the $n(314 \text{ nm})$ variation can serve as an indicator of the degree of curing in the dielectric thin film. The reduction in the $n(314 \text{ nm})$ value decreased as the temperature or time approached the high end of the curing process window, indicating that the curing process was nearly completed. After determining the correlation between curing parameters and optical constants, a single-parameter empirical interpolation model could be employed.



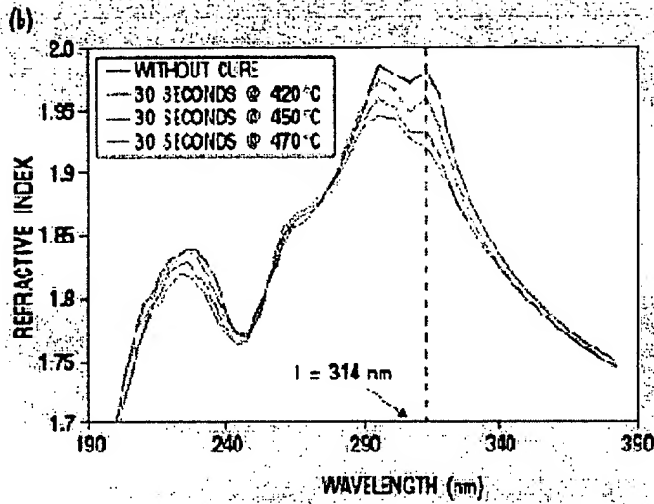


Figure 1: VASE measurements of the ultraviolet RIs of low-k resins cured under different conditions: (a) cure-time dependence; (b) cure-temperature dependence.

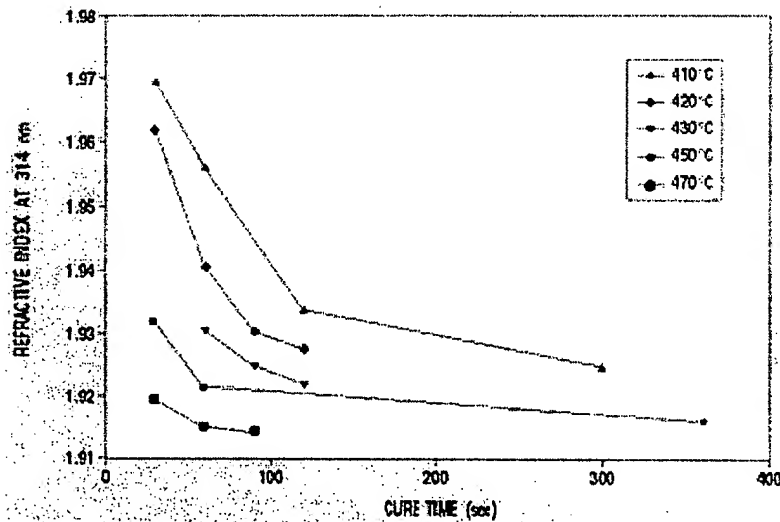


Figure 2: VASE measurements of RIs at 314 nm at different curing times and temperatures.

In a process-sensitive thin film, process parameter variations can lead to changes in the film's $n(\lambda)$ and $k(\lambda)$ values. That is the case for the cured resins investigated in this study. Monitoring the degree of curing in these dielectric thin films on an automated metrology tool requires a parametric dispersion model for the optical constants of these resins, which represents the functional dependence of $n(\lambda)$ and $k(\lambda)$ on λ in terms of a small number of adjustable parameters. This model should be able to describe the $n(\lambda)$ and $k(\lambda)$ of these resins cured under all conditions within the whole process window.

An empirical interpolation model is an effective single-parameter dispersion model whose adjustable parameter normally corresponds to a process-dependent physical property (e.g., the crystallinity of polysilicon or the alloy fraction ratio of SiGe).⁹ It consists of a database of optical constant spectra for a given material. Each spectrum corresponds to a specific process condition. When fitting reflectance or

ellipsometry data of a sample prepared under unknown conditions (within the process window), the sample optical constants are interpolated between known spectra in the database. With only one adjustable parameter, the correlation between thickness and optical constants can be minimized during data analysis.

To develop an empirical interpolation model for these low-k resins, optical constant spectra of five samples cured under different conditions were chosen. The $n(314\text{ nm})$ values of these samples, which are presented in Table I, were evenly distributed between the lowest and highest values in the process window. The term *cure index* was used to refer to the adjustable parameter in the model. A cure index value of 0 was assigned to the uncured sample, which had the highest $n(314\text{ nm})$ value, while an index of 1 was assigned to the cured sample with the lowest $n(314\text{ nm})$ value. The other three cured samples were evenly spaced between the samples with the highest and lowest $n(314\text{ nm})$ values. The cure indices were determined by proportioning the $n(314\text{ nm})$ values of the five samples between 0 and 1 as follows:

$$\text{Cure index} = [n(314\text{ nm}) - 1.9791] / [1.9161 - 1.9791]$$

Automated Production Metrology

The empirical interpolation model for low-k resins was tested on the automated thin-film metrology tool. Using the tool's reflectometer mode, both film thickness and optical constants were determined from a recipe set up to measure absolute UV-visible reflectance.¹⁰ Figure 3, which presents a reflectance scan and fit result based on the empirical interpolation model, shows an excellent match between the experiment and the model. To become production worthy, however, the metrology method must demonstrate high levels of precision, stability, accuracy, sensitivity, and throughput.

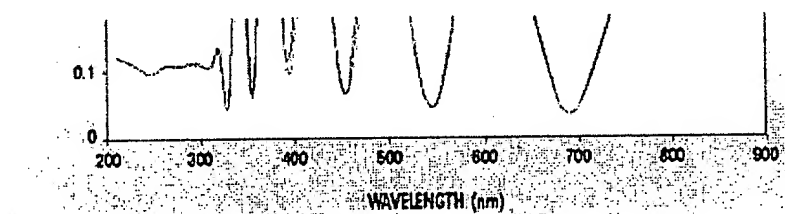


Figure 3: Fit result of reflectance data from low-k resin treated at 400°C for 300 seconds shows an excellent match between the experiment and the model.

| Sample | Cure | $n(314\text{ nm})$ | Cure |
|--------|------|--------------------|------|
|--------|------|--------------------|------|

| No. | Condition | | Index |
|-----|---------------------|--------|-------|
| 1 | Without cure | 1.9791 | 0.00 |
| 2 | 30 seconds @ 410°C | 1.9694 | 0.15 |
| 3 | 60 seconds @ 400°C | 1.9513 | 0.44 |
| 4 | 60 seconds @ 430°C | 1.9307 | 0.77 |
| 5 | 360 seconds @ 450°C | 1.9161 | 1.00 |

Table I: Data for five samples whose optical constant spectra made up the single-parameter interpolation model.

The measurement precision and dynamic stability of the metrology tool were tested by examining several wafer samples that had varying film thicknesses or had been cured under different conditions. One of the worst cases was a 1000-Å-thick resin sample that was loaded 10 times and measured in five places at each load (one at the center and four around the edge). The results revealed that the standard deviation of $n(314 \text{ nm})$ for 10 loads at each site was <0.001 and that the variation of the average value of $n(314 \text{ nm})$ for five sites was also <0.001 from load to load. Since the uncertainty was $<2\%$ of the total range of the variation in $n(314 \text{ nm})$ that may be induced by curing (about 0.063), it was concluded that this measurement technique provides sufficient precision and stability to monitor the curing process of these low-k resins. Table II lists the statistical results of the average thickness and $n(314 \text{ nm})$ of each load.

| | Mean | Standard Deviation |
|---------------------|--------|--------------------|
| Thickness (Å) | 1064.8 | 1.14 |
| $n(314 \text{ nm})$ | 1.929 | <0.001 |

Table II: Statistical test results from a worst-case 1000-Å-thick sample that was loaded 10 times and measured at five places after each load.

By comparing $n(314 \text{ nm})$ values measured by the automated in-line tool to the benchmark values from the VASE measurement, measurement accuracy could be evaluated. Figure 4 shows that the data points fit a straight line with a slope of 0.9788 and a y-intercept of 0.0441, indicating excellent correlation between the two types of measurements. This close match demonstrates that the refractive index measurements using the reflectometry mode of the in-line tool were accurate for monitoring the curing process.

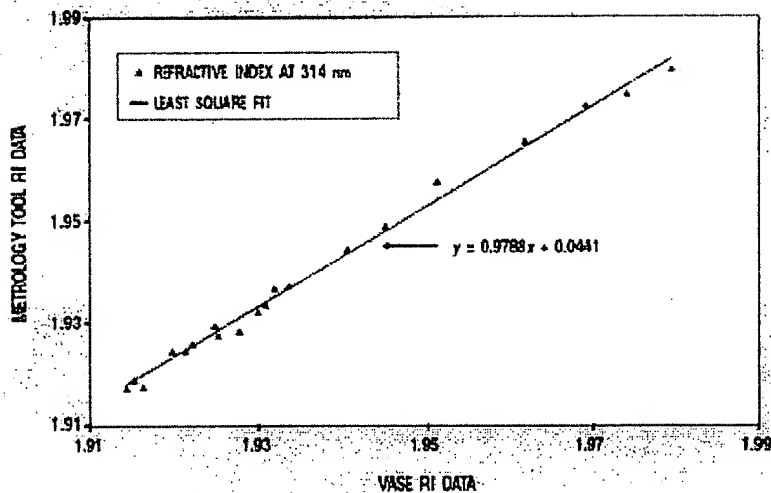


Figure 4: Correlation between RI at 314 nm measured by the VASE and the automated metrology tool shows an excellent match between the two types of measurements.

The model's sensitivity to the curing process as well as the feasibility of determining the film thickness and optical constants simultaneously are seen in Figure 5, which plots three model-predicted reflectance spectra corresponding to three cure conditions: 30 seconds at 410°C, 30 seconds at 420°C, and 60 seconds at 420°C. All three spectra were generated based on the same thickness value of 7400 Å. The difference between the spectra is strictly a result of the difference in the degree of cure. Figure 5 clearly demonstrates that there is significant difference among these spectra in the UV spectral region (between 310 and 400 nm), which can be readily measured by a reflectometer. Film thickness information, on the other hand, can be found in the period of the oscillating portion of the reflectance spectrum. By fitting a broadband UV-visible reflectance spectrum, film thickness can be measured quickly and reliably, and the cure-induced change in the resins can be monitored.

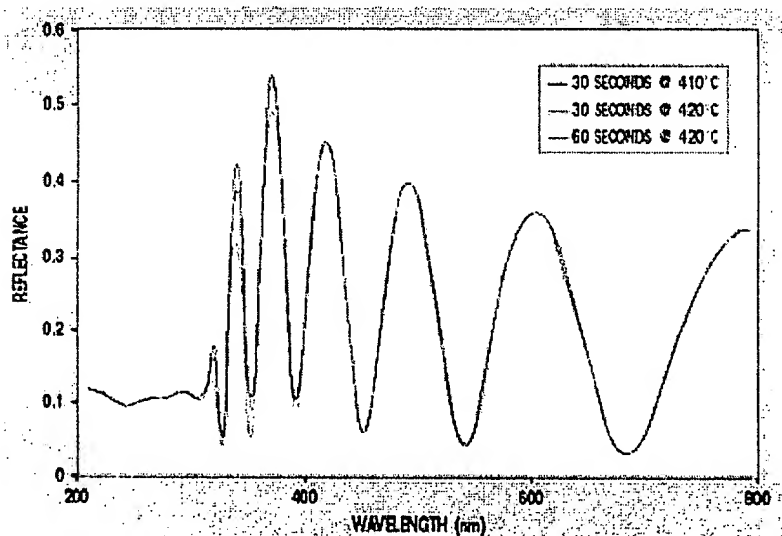


Figure 5: Reflectance sensitivity predicted by the empirical interpolation model for low-k resins under

three different cure conditions; the difference between the spectra is a result of the difference in the degree of cure.

Measurement throughput depends on the speed of reflectometer data acquisition and analysis, as well as the focusing speed of the sample stage (e.g., the system's robotics). To boost the UV signal-to-noise ratio, a broadband measurement requires longer detector integration time than a visible reflectance measurement, leading to decreased throughput. The metrology tool used in this study performed measurements at five sites on an 8-in. wafer in approximately 1 minute with focusing on each site. New-generation spectrophotometer heads, robotics, and data-handling software are in the final testing stages, promising much improvement in measurement throughput.

Conclusion

In order to serve as adequate low-k ILD materials, organic spin-on polymers must be properly cured. In-line characterization of the curing process is crucial because it improves equipment efficiency, eliminates the need for monitor wafers, and reduces wafer scrap and wasted work. The curing process of low-k resins can be monitored with broadband reflectometry, and film thickness and the degree of curing can be calculated simultaneously. Because of its fine measurement spot size and pattern recognition capability, this technique can be used to monitor the curing process of low-k thin films on product wafers. The method described here for monitoring SiLK should be applicable to other low-k materials if the change in the optical constants induced by polymerization is monotonically related to the process parameter and the metrology tool is sensitive enough to measure the change in optical constants. Additional studies must be performed to determine the feasibility of performing metrology measurements on product wafers with device structures and the efficacy of using integrated in-line metrology tools.

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Exhibit F

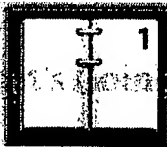

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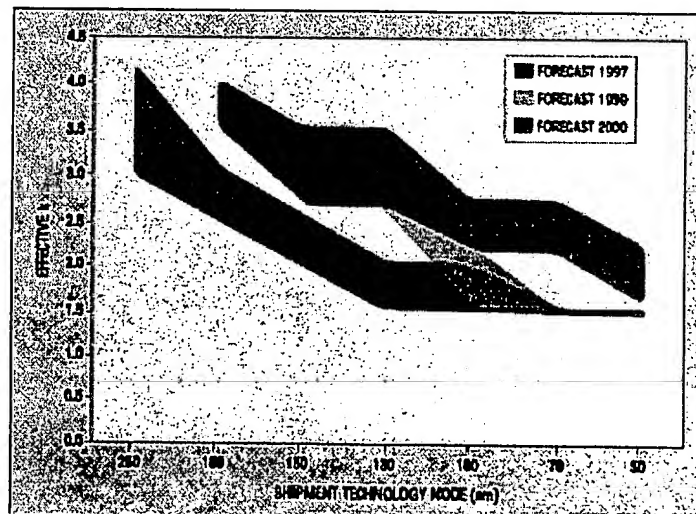
Media Kit

Addressing the challenges of spin-on low-k dielectric dispense management

Josh H. Golden, J. Eric Carrubba, and Jay Jung, *Microbar*

Optimizing dispense tool designs and SOD management will require cooperation between equipment manufacturers, materials suppliers, and end-users.

During the last five to eight years, the semiconductor industry, academia, and government labs have been engaged in an intense worldwide research effort in preparation for the integration of low-k dielectric materials in ICs at the 180-nm technology node, which was scheduled for first shipments in 1999. This effort was driven by the desire to uphold Moore's Law and the various revisions of the industry roadmap.¹⁻⁴ However, the transition from silicon dioxide films applied by chemical vapor deposition (CVD), which have a dielectric constant (k) value of 4.1 to 4.2, to inorganic, organic, and hybrid materials with k values of less than 3.0 has proven to be more difficult than predicted. Figure 1 illustrates the delayed adoption of low-k dielectrics based on the roadmaps.^{4,5}



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Figure 1: The delayed implementation of low-k materials at each device generation. (Adapted from reference 5.)

As both research and delays continue, a debate has been raging over the choice of techniques for applying low-k materials. Both spin-on and CVD candidates present integration challenges because their properties differ from those of the benchmark, SiO_2 , which is hard, relatively inert, thermally stable, and easy to deposit or grow.⁶⁻⁸

Table I compares the dielectric properties of SiO_2 to those of organic spin-on dielectrics (SODs) and $\text{Si}_w\text{C}_x\text{O}_y\text{H}_z$ CVD films. The organic SODs may present a greater integration challenge than do the hybrid $\text{Si}_w\text{C}_x\text{O}_y\text{H}_z$ CVD films, which are more "silica-like." For example, SiO_2 has a modulus of 72 GPa, while a leading polyarylene SOD exhibits a modulus of 2.7 GPa.⁸ This approximately 27-fold difference in modulus has a significant effect on the efficacy of chemical mechanical planarization (CMP) processes, which subject copper interconnect layers to significant shear forces.⁹

| Property | Material | | |
|-------------------------|-----------------|---|--|
| | Silicon Dioxide | Organic SODs | CVD $\text{Si}_w\text{C}_x\text{O}_y\text{H}_z$ |
| Dielectric Constant (k) | 3.9—4.5 | 2.4—2.85 | 2.5—3.2 |
| Modulus (GPa) | 72 | 2—3 | 4.2—10.8 |
| Hardness (GPa) | 8.7 | ~0.25 | 0.23—1.41 |
| Composition | SiO_2 | Various: Poly(arylene ether) Poly(arylene) Poly(benzoxazole) | 15—22% Si, 17—23% C, 18—30% O, 26—45% H |

Table I: Comparison of some properties of silicon dioxide with those of SODs and carbon-doped siloxane film. (Adapted from reference 8.)

In addition to the modulus, heat dissipation is another important issue with low-k candidates, because their thermal conductivity is typically 20 to 30% lower than that of SiO_2 .^{6,7,10} Poor heat dissipation can cause localized metal-line heating, which eventually leads to device failure. Thermally induced stresses within the low-k film may lead to delamination because of mismatches between the film's coefficient of thermal expansion (CTE) and the substrate CTE. For example, silicon dioxide's CTE is 0.5 ppm/°C, aluminum's CTE is 23.1 ppm/°C, and copper's CTE is 16.5 ppm/°C. Because the CTE of many organic dielectrics is >50 ppm/°C, CTE mismatches can lead to high tensile stresses following thermal cycles.⁷

A successful low-k candidate must display several critical material properties: chemical resistance to oxidation and moisture absorption during plasma ashing, stripping and cleaning, and CMP processes; thermal stability (no weight loss or shrinkage following repeated

thermal stability (no weight loss or shrinkage following repeated isothermal soaks at 400°C); and the ability to adhere to substrates, including liners and barriers, in order to withstand the shearing and delamination forces exerted by the CMP process.

A variety of options have recently emerged to address some of the challenges associated with using low-k materials in ICs. These include delaying the implementation of such materials as well as new integration schemes. Approaches include:

- Combining new circuit architectures with aluminum and low-k dielectrics to achieve lower resistance-capacitance (RC) delays.^{2,3,5,11}
- A fluorinated silica glass technique.
- An embedded approach that uses low-modulus and low-thermal-conductivity low-k materials only between the lines, where 80% of the reduced RC-delay benefit is realized. At the via level, SiO₂ or a spin-on glass (SOG) technique may be used for added strength and improved thermal conductivity.^{5,12}
- Using a polyarylene thermoset at the lower five tight-pitch copper levels and fluorinated silica glass at the top three global wiring levels for strength and heat dissipation.^{8,13}
- Using relatively inert ($k \sim 4.5$) silicon carbide as a CMP barrier film and hard mask to protect a potentially reactive low-k film.⁶

While these possibilities raise many interesting issues, this article focuses on some of the benefits and concerns associated with the adoption of SODs. The need for dispensing equipment especially designed for this application and the need for SOD material management capabilities are particularly emphasized.

SODs versus Hybrid CVD Films

The April 2000 announcement by IBM that it will adopt Dow Chemical's SiLK resin as an insulator for its dual-damascene copper processes at the 130-nm technology node represented the first major breakthrough in the spin-on versus CVD debate.¹³ Since then, UMC in Hsinchu, Taiwan, and Altis Semiconductor (an IBM-Infineon joint venture) in Corbeil Essones, France, have announced that they will also use the 130-nm process technology developed with IBM, and it is likely that more companies will announce their adoption of SiLK sometime this year. However, other 2.2–3.1-k SODs are still being considered, including Honeywell's HOSP, a hybrid siloxane-organic polymer; Dow-Corning's XLK, a porous hydrogen silsesquioxane (HSQ); Schumacher's MesoELK, a porous silica product; and JSR's MSQ hybrid.

SODs' main competitors at the 130-nm technology node are hybrid Si_wC_xO_yH_z CVD films with dielectric constants in the 2.5–3.0 range.

These films are more like SiO_2 than the SODs are, and they have a lower density and lower polarizability because of their pendant organic methyl groups. Other advantages of the films include the good availability and low cost of the precursor materials, as well as their ability to be processed with existing CVD tool sets.^{5,14,15} However, the $\text{Si}_w\text{C}_x\text{O}_y\text{H}_z$ films are relatively soft compared to SiO_2 (0.23–1.41 versus 8.7 GPa).⁸ It also appears that $\text{Si}_w\text{C}_x\text{O}_y\text{H}_z$ hybrids with a dielectric constant of <2.5 display film properties that will be inadequate for the 100-nm technology node, although research efforts are under way by materials and CVD tool suppliers to address such issues.

On the other hand, SODs have certain intrinsic material advantages over low-k CVD films. For example, a SOD material typically has a well-defined chemical composition that undergoes a specific and well-characterized physicochemical change (i.e., cross-linking) during thermal cure. In contrast, the composition of $\text{Si}_w\text{C}_x\text{O}_y\text{H}_z$ films may vary from recipe to recipe and from tool to tool, as a CVD process is developed and qualified.

In addition, because SOD compositions are created using established chemical synthesis and modification procedures, the preparation of porous materials with dielectric constants as low as 2.0 is readily achievable. A variety of approaches are being used by synthetic chemists in the preparation of both open- and closed-cell porous networks with optimized pore sizes and distributions. These include the use of porogens, such as a high-boiling-point solvent (Dow-Corning's XLK), or a thermally decomposable organic molecule, such as a dendrimer.¹⁶

Mesomorphous silicas represent another class of SODs. To create these materials, an organized porous silica or $\text{Si}_w\text{C}_x\text{O}_y\text{H}_z$ network is formed by templating silicon-containing precursors with organic surfactants in a mixed-solvent system. Sol-gel or sol-gel-like ordered structures are then created in a prebake step, followed by the burning off of the templating organic molecules. In contrast to these porous-SOD technologies, the preparation of porous materials for plasma deposition is a greater challenge because it is more difficult to control pore size, achieve pore-size homogeneity, and create closed-cell architectures. Both CVD equipment suppliers and materials manufacturers are researching ways to meet this challenge.

Another advantage of SODs over films applied by CVD is that the sensitivity of SODs to moisture, oxygen, and heat is generally understood and somewhat predictable. Thus, necessary precautions can be integrated into handling and dispensing procedures without extensive experimentation. The ease with which the handling and dispensing of these materials can be optimized, and the relative

simplicity of the spin-on process, help to hold down the end-user's cost of ownership (COO). In contrast to SODs, the optimization of $\text{Si}_w\text{C}_x\text{O}_y\text{H}_z$ processes requires rigorous experimental designs in order to achieve the best combination of mass flow and multiple-component precursor ratios, as well as the necessary process tool parameters. Significant effort must therefore be expended to maintain the process window, the cost of which typically has not been included in COO studies.¹⁵ Another hidden cost is that incurred in the transference of the process from one site or tool to another, which can be difficult and time-consuming.

Proponents on both sides of the dielectric debate continue to argue the respective strengths and weaknesses of the various technologies, from materials properties to ease of use to COO.¹⁵ However, it appears that a combination of SOGs, SODs, and CVD films will be used in conjunction with new integration schemes at both the 180- and 130-nm technology nodes, and possibly at the 100-nm node as well.

SOD Dispense Tool Design Issues

The implementation of SODs will require that new cooperative relationships be developed between tool manufacturers, materials suppliers, and end-users. Formerly, when spin-on tools were used only for photoresist, there was a mostly closed loop between spin-on track manufacturers and dispense cabinet providers. Now, however, because of a variety of factors, including the existence of multiple SOD candidates with different chemistries and their high costs (>\$1000/L), it has become critical for equipment manufacturers, materials suppliers, and end-users to work closely to ensure that the chosen SOD will be optimally dispensed onto the wafer.

There are significant differences between photoresist dispense and SOD dispense that can affect tool design and process parameters. After dispense, a photoresist undergoes a variety of processes, including prebake, exposure, development, and cure, after which it is stripped off the wafer. But because an SOD is integrated permanently into the device structure, careful attention must be paid to preventing contamination and maintaining the correct material thickness and composition during the dispense process. Failure to control particles and other contaminants results in the classic "fly stuck in amber" scenario, which can ultimately lead to device failures.

In addition, while most photoresists can be handled similarly, each SOD has a different set of handling criteria because of its unique chemistry and physical properties. Dispense tool manufacturers must understand these properties and those of the carrier solvent(s), applying this knowledge to the design of their hardware and software. Knowledge of the SODs' properties must drive the choice of tool component materials and the integration of safeguards against

contamination. Table II lists some SODs and relevant information on their properties.

| Property | SOD Material | | |
|---|--|-----------------------------------|--------------------------|
| | Dielectric A | Dielectric B | Dielectric C |
| Dielectric constant (k) | 2.65 | 2.0–2.5 | 2.5 |
| Type (solute) | Polyarylene | Porous HSQ | Organic siloxane polymer |
| Solvent(s) | Butyrolactone, cyclohexanone, mesitylene | 2-Pentanone, toluene, tetradecane | Propyl acetate |
| Solute moisture sensitivity (25°C storage, 1 week) | No | No | Yes |
| Hygroscopic solvent(s) | Yes | Yes | Yes |
| Thermal sensitivity (25°C storage, 1 month) | No | Yes | Yes |
| Oxygen sensitivity (25°C storage, 1 week) | No | No | No |
| Particle formation potential (25°C storage, 1 week) | No | No | Yes |
| Flammable mixture | Yes | Yes | Yes |

Table II: Comparative data on SOD properties that are useful in the design and operation of dispense tools. (Data collected from materials suppliers' material safety data sheets and personal communications.)

Tool manufacturers must also consider the properties of SOGs and their derivatives, which may be used as etch stops and barriers in conjunction with a low-k SOD.^{6,11} Because the wetting characteristics of SOGs typically are optimized for gap-fill and planarization applications, these materials are prone to wicking and may form small inclusions in nooks and crannies in the dispense apparatus' fluid path. This phenomenon can lead to the formation of cross-linked silica-like particles upon solvent removal or evaporation. One solution to this problem is to use low-roughness-average materials for the dispense tool's fluid paths and to provide a solvent-saturated inert-gas environment within the tool.

Many polar carrier solvents are hygroscopic or may undergo acid or base hydrolysis under certain conditions. While gross solvent hydrolysis is unlikely in a well-engineered dispense apparatus, even small amounts of moisture can have a deleterious effect on the composition of SODs that contain metastable siloxane oligomers or

composition of SODs that contain metastable siloxane oligomers or silane or siloxane adhesion promoters. Therefore, careful attention to SOD bottle changes and the use of an inert-gas backfill to prevent oxygen or moisture infiltration is required.

Some SOD materials are thermally sensitive and thus require a temperature-controlled environment to prevent spontaneous polymerization, cross-linking, or condensation of oligomeric moieties, which can result in particle formation. Possible measures to prevent such reactions and to optimize dispense temperature include transferring the cold-preserved SOD from the container to a secondary, temperature-controlled reservoir and the inclusion of thermal controls ($\pm 1^\circ\text{C}$) along the dispense tool's fluid path. Temperature control is important not only for preventing particle formation, but also for helping maintain the fluid viscosity necessary for correct on-wafer film thickness.

In addition to temperature controls, an effective SOD dispensing system should also provide solvent-saturated backfill and automated flush-purge capabilities for preventing particle formation and deposition within the dispense apparatus. While solvent-saturated backfill ensures a wetted path for the SOD with a like solvent or solvent mixture, an automatic solvent flushing protocol cleans the system and then dries it with an inert gas. An automated flush-purge system flushes and cleans dispense lines between batch changes and helps perform routine maintenance. The ability to isolate and purge portions of the fluid path for cleaning and servicing in a multitrack dispense tool while processing continues uninterrupted will become increasingly important as SOD end-users ramp up capacity and move from using 1-L containers to bulk delivery systems.

Another challenge facing SOD dispense systems is the removal of particles and microbubbles, which is accomplished through one-pass filtration or filtration with recirculation, depending on the material's requirements. A study of particle and microbubble removal using filtration with recirculation was conducted at the Honeywell Electronic Materials Star Center (Sunnyvale, CA) using a PMS M65 particle detector from Particle Measuring Systems (Boulder, CO) integrated with the Low-k 2.0 SOD/SOG dispense system from Microbar (Sunnyvale, CA).

In the study, the particle-size detection limit was $0.065\ \mu\text{m}$, and the dispense tool included a $0.065\text{-}\mu\text{m}$ filter. Data were collected at 25.5°C at a sampling flow rate of $0.6\ \text{ml/min}$. Figure 2 shows the results of this study for an $\text{Si}_w\text{C}_x\text{O}_y\text{H}_z$ hybrid SOD that is sensitive to both temperature and moisture and thus has the potential to form insoluble particles. As the figure indicates, acceptable particle reduction was achieved after approximately 3 hours, or 175 recirculation cycles. While it proved difficult to quantitatively differentiate microbubbles from particles because of their similar size and an inability to isolate each component, the study found that both particles and microbubbles

each component, the study found that both particles and microbubbles were removed to acceptable levels without affecting film thickness or quality.

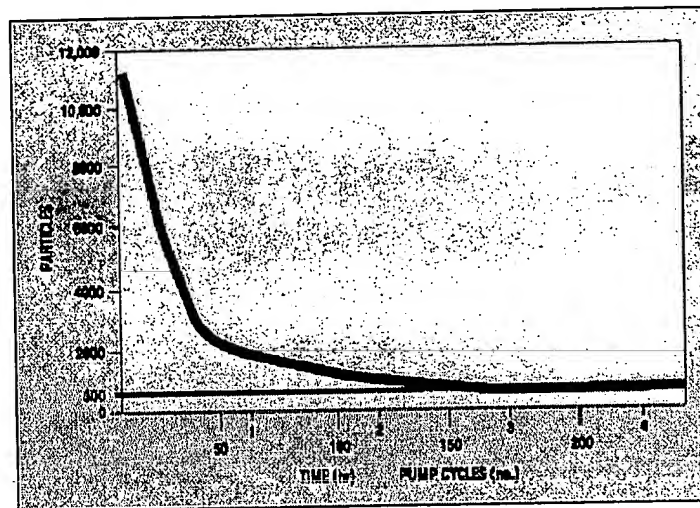


Figure 2: Data collected for an $\text{Si}_w\text{C}_x\text{O}_y\text{H}_z$ SOD hybrid using a particle detector that was integrated with an SOD dispense system equipped for filtration with recirculation.

Filter selection can be critical to achieving good results. While the choice of filter material and type is a materials-compatibility issue, the choice of pore size can directly affect SOD composition and molecular weight. Many SODs consist of oligomers or prepolymers and/or polymeric materials that have a specific molecular-weight distribution and average thermodynamic radius. A filter whose pore size is too small has the potential to fractionate this molecular-weight distribution and thus remove a certain population of oligomers or polymers from the SOD mixture. This can adversely affect the material's on-wafer thickness, physical properties, and, possibly, device yield performance. Because the molecular-weight distribution may also be affected by shearing forces (polymer chain cleavage), careful fluid-path design and selection of transfer pumps and associated apparatus are necessary for some SOD materials as well.

Dispense tool manufacturers also must address environmental safety and health concerns, including the prevention of spills and leaks of reactive, noxious, and flammable spin-on materials. Double-containment piping, spill sensors, quick-release fittings, and proper ventilation of and safe access to the dispense cabinet are among the features that can be included. Multiple-track SOD dispense schemes with built-in safety features can also increase tool efficiency and process uptime. Safety features that facilitate bottle change will also prove beneficial as SOD use increases.

SOD Management

The tracking and management of SODs is another important reason why materials suppliers, dispense tool manufacturers, and their

may maintain suppliers, dispense tool manufacturers, and their customers must develop cooperative relationships. Issues that affect an SOD's life span include supplier batch and materials information, the transport environment from the supplier to the end-user, and the dispense protocol and associated parameters. A combination of hardware and software strategies can be used to compile, manage, and share the required data. For example, SOD containers can include a wireless tracking device that operates in the RF spectrum and is located in a well or integrated in a dip tube. Activated at the SOD point of fill, the device logs such transport parameters as time, thermal history, shock, and tilt. Because some SODs are sensitive to temperature and shock, monitoring these parameters can pinpoint potential upsets that can lead to the spoilage of the material.

Batch-tracking capability is important for maintaining quality control, process stability, inventories, and orders. In a comprehensive SOD-tracking scheme, data collected at the point of fill, during shipment, at fab delivery, and before bottle installation in the dispense tool can be uploaded into a software package that allows analysis and the sharing of data among authorized users. Analytical features can include troubleshooting and go/no-go protocols to be implemented when an SOD shipment enters the fab receiving area and again during bottle changeover at the dispense tool. Wireless tracking can also be integrated with bar code schemes to enable analysis. Figure 3 illustrates a possible combination of hardware and software technologies that can serve as the anchors of an integrated system for the management of SODs.

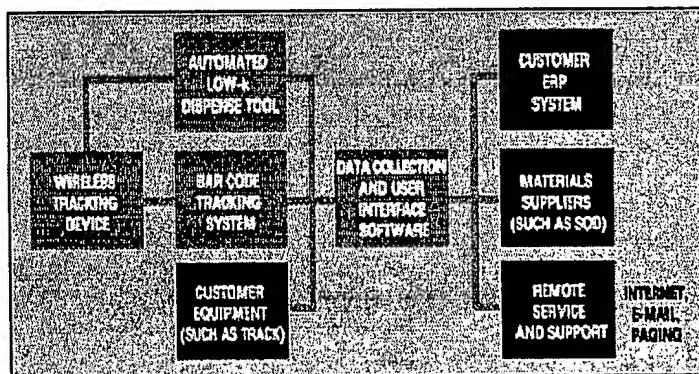


Figure 3: Schematic showing how a central software chemical-management package can link customers, materials suppliers, and dispense tool manufacturers with data from monitoring devices and process tools.

One example of such a system, developed by Microbar, includes the company's software platform and wireless tracking device. The software is capable of collecting data from the manufacturer's tool sets and those of other equipment suppliers, and can transmit both data and analytical results to materials suppliers and end-users via interfaces and the Internet. The types of information provided by the platform include histograms of equipment uptime and performance, maintenance and service logs, security information to be used by

process engineers to validate chemicals, and mass balances of chemical-distribution loops for environmental and operational expense measurements. The implementation of such a system should minimize equipment COO by promoting the efficient utilization of expensive SOD materials.

Conclusion

Although the debate over the optimal technique for applying low-k dielectrics continues, some semiconductor manufacturers have announced that they are adopting SODs. Such materials have several advantages over CVD films, including their well-defined chemical composition and physical characteristics. However, their adoption will require new cooperative relationships between equipment manufacturers, materials suppliers, and end-users. Tool designers, in particular, must understand the properties of the many available SOD materials to ensure that they are dispensed optimally. Dispense tools must be designed to enable the prevention and removal of contamination, and must have built-in safety features. SOD management schemes that link suppliers and customers will also be needed to minimize cost of ownership and facilitate the efficient use of the costly materials.

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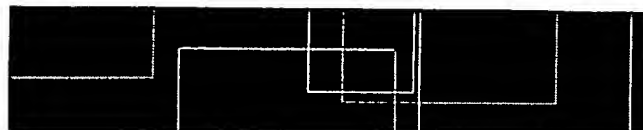
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Spin-on-Glass

Product Description:

ACCUGLASS T-11 Spin-on-Glass (SOG) Series is a family of methylsiloxanes that combine organic groups on an inorganic polymer backbone. It is used in production by IC manufacturers for interlevel and overcoat passivation smoothing. This product was designed for microelectronic applications requiring a high quality dielectric film of up to 3,500 Å, with a κ value of 3.8.

ACCUGLASS T-12B Spin-on-Glass (SOG) Series is a family of advanced spin-on dielectrics for use in the fabrication of integrated circuits. Chemically, this product belongs to the methylsiloxane family of polymers. This crack-resistant material features one-half micron single coat thicknesses, low shrinkage upon cure, and a κ value of 3.1.

ACCUGLASS T-14 Spin-on-Glass (SOG) series is a family of siloxane polymers formulated to fill narrow, high aspect ratio spaces without voiding and to more completely planarize multi-level metal devices. The organic content of the material is approximately 10%. With a κ value of 3.8, this material has good adhesion to silicon, thermal oxide, CVD oxide and aluminum. This polymer is formulated for microelectronic applications to better planarize all ICs, to fill narrow gaps in 0.5 μm to 0.35 μm devices and to extend the use of SOG for next generation ICs.

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Cover photo: Electron micrograph of contact holes filled with CVD tungsten plugs (see Chapter 8). The diameter is 0.25 micron. Courtesy of the National Nano Device Laboratories. National Science Council, R.O.C.

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The characteristics of O_3 -TEOS can be summarized: (1) It can be deposited by APCVD or SACVD at 250 to 400°C at rates depending on both temperature and O_3 concentration; (2) the oxide has good step coverage and can fill higher aspect ratio gaps than PECVD TEOS; (3) the film properties are sensitive to surface conditions; and (4) the film properties are functions of deposition temperature and O_3 concentration. Higher temperature and higher O_3 concentration lead to higher-quality oxides but at the cost of lower deposition rates.

O_3 -TEOS oxide is somewhat porous, with a shrinkage of 1 to 3% after a 450°C anneal. When left in air, the film absorbs moisture gradually. The water absorption can cause gate oxide damage and via poisoning and has to be avoided. Figure 27 shows the process sequence of a structure where the O_3 -TEOS oxide is etched back to provide a gap filler. In this approach, a thin layer of O_3 -TEOS oxide is first deposited to fill the gap (Fig. 27a). If it is left on top of the metal, then moisture absorption and re-release during metal deposition will lead to oxidation and poisoning of the metal contact (Fig. 27b). Instead, an RIE process is used to remove the O_3 -TEOS oxide (Fig. 27c). Finally, a thick PECVD TEOS is deposited to complete the ILD process (Fig. 27d). This process can fill narrow gaps with aspect ratios of about 1.3 to 1.5. For higher aspect ratio gaps new deposition processes are needed.

Spin-on-glass (SOG)

Spin-on-glasses are widely used for low-cost fabrication of IC circuits. There are two basic types of SOG: organic and inorganic. The inorganic SOGs are silicate-based, and the molecular structure is shown in Fig. 28a. The structure of an organic

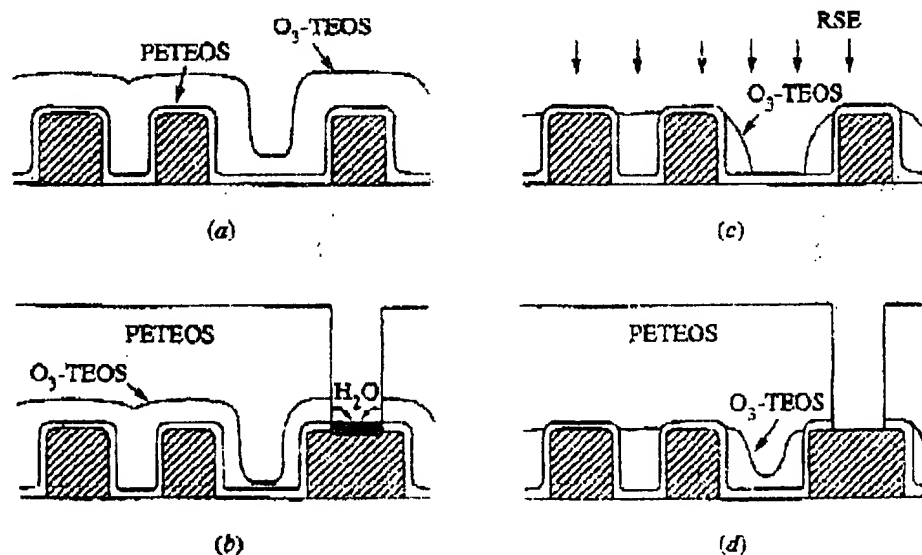
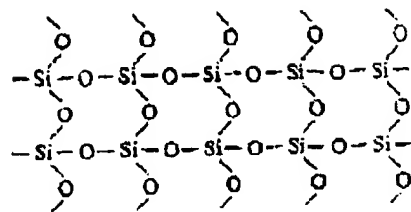


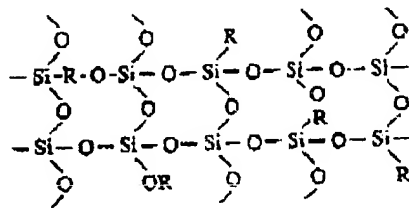
FIGURE 27

O_3 -TEOS/PECVD ILD processes: (a) PECVD liner before O_3 -TEOS deposition; (b) if the O_3 -TEOS is left over metal, the exposed portion may absorb moisture and outgas, oxidizing the metal contact; (c) to avoid contact problems, remove O_3 -TEOS from contact area by RIE; (d) a process that leaves no exposed O_3 -TEOS.

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(a)



(b)

FIGURE 28

Schematic of the molecular structures of SOGs:
(a) silicate SOG; (b) siloxane SOG.

SOG is shown in Fig. 28b. The organic, or siloxane-based, SOGs are featured with radical groups replacing or attaching to oxygen atoms. From these two basic structures, numerous commercial modifications have been made to adjust the molecular weight, the viscosity, and the final film properties for specific applications.

After full curing, the silicate SOG behaves like SiO_2 . It does not absorb water in significant quantity and is thermally stable. However, the volume shrinkage during curing is large, and as a result the silicate SOG develops high stress (~ 400 MPa) and cracks easily during curing. Applying only a thin layer (100 to 200 nm) avoids the cracking problem. To build up thicker layers to fill gaps, multiple application and curing are needed. The ability to fill narrow gaps is also limited since low-viscosity SOGs (which have better filling properties) usually have higher shrinkage.

Silicate SOGs can be modified by adding P (up to 4 at%) to produce a softer SOG. The P is incorporated into the silicate by replacing Si atoms and breaking Si-O bonds. Therefore, adding P to a silicate SOG creates a structure similar to the siloxane SOG (Fig. 28b). The modified SOG (called P-silicate) is softer and has less shrinkage. The stress after curing is reduced to ~ 200 MPa, and the SOG is more crack-resistant. It also has a better gap-filling capability and can be applied in thicker layers. Adding P, however, increases the water absorption and the modified SOG is both mechanically and electrically less stable than the silicate SOG.

Siloxane SOG, on the other hand, retains carbon even after curing. It has lower stress (~ 150 MPa) and can be applied in thicker layers. The organic SOG, however, is sensitive to water absorption, cannot tolerate even moderate temperature ($> 400^\circ\text{C}$) cycles, and has poor mechanical strength. Organic SOGs may also become unstable under plasma exposure. The amount of organic component in the siloxane is an adjustable parameter, and the SOG can be tailored to specific applications.

Because silicate SOGs are more SiO_2 -like, they are used mainly for gap filling even when multiple applications and curing are necessary. The organic SOGs are more suitable for sacrificial SOG, e.g., the planarizing layer for etchback.¹³⁹ (See

Section 8.5.3.) Because of water absorption and subsequent via poisoning by SOGs, they are used only for gap fill and the excess is removed. The processing sequence shown in Fig. 27a, c, d applies to SOG with the simple replacement of SOG for O_2 -TEOS oxide.

Because both the material and the method of application are inexpensive, SOG now becoming a popular low-cost ILD for gap fill. The etchback step (Fig. 27c) is expensive and a potential source of particles. Therefore, a nonetchback SOG process is important. Recent work has reported spin-on materials that can be left on metal without jeopardizing via contacts.^{140,141} Fluorine treatment of a siloxane SOG to use as a nonetchback ILD is also reported.¹⁴² No large-scale testing has been completed for these new materials and therefore the acceptance is still experimental. Although some fully cured SOGs have lower dielectric constants (3.0 to 3.6 compared with 3.9 to 4.5 for SiO_2) than oxides, in general SOGs absorb water and both the dielectric and mechanical properties are inferior to those of PECVD oxides. Lower cost, good gap-filling capability, and better ILD planarity still make SOG an attractive alternative to PECVD oxide. Currently, SOGs are used for manufacturing devices from $> 1 \mu m$ to $0.35 \mu m$ in size. *

Bias-sputtered quartz

SiO_2 can be deposited by rf sputtering from a quartz target at room temperature. The sputter-deposited oxide is porous, has poor step coverage, and is not suitable for ILD use. Adding a dc bias, however, changes both the film quality and the step coverage drastically. The ion bombardment during sputter deposition provides sufficient energy to densify the oxide to a level similar to thermal oxide. The energetic ions sputter off the overhang from feature edges and create shadow-free deposition. Consequently, a complete filling of gaps between metal lines is achieved. Figure 29a shows the resputtering rate of SiO_2 as a function of ion incident angle.¹⁴³ The sputtering rate is low at normal incident angle and stays low up to about 30° . It has a broad peak at about 70° and then falls off sharply to zero at 90° . In bias quartz sputtering the deposition rate and bias are adjusted so that the deposition rate is between the minimum and the peak of the resputtering rates. If the deposition rate falls below the minimum resputtering rate, then the substrate is eroded and there is no deposition. If the deposition rate is higher than the peak of the resputtering rate, then bias sputtering is not effective, resulting in poor step coverage. Figure 29b shows the layer profiles over different features for biased sputter deposition. Note the building of sharp ridges over metal features. As the SiO_2 grows in thickness the ridges on smaller features disappear first. Bias-sputtered deposition (Fig. 29b) can fill narrow, high-aspect ratio gaps solidly when parameters are properly adjusted.

Bias-sputtered quartz has not been used widely for ILD applications for the following reasons: (1) the low deposition rate; (2) particulates from chamber walls and shielding; (3) metal contamination from chamber parts; (4) radiation damage from energetic electrons (which produce x-rays); and (5) high compressive stress (~ 1000 MPa) of the film, which cracks easily. However, the benefits of bias sputtering during deposition have been recognized. The combination of bias sputtering with plasma-enhanced CVD of SiO_2 led to the most promising ILD process (see next section on HDP oxide).

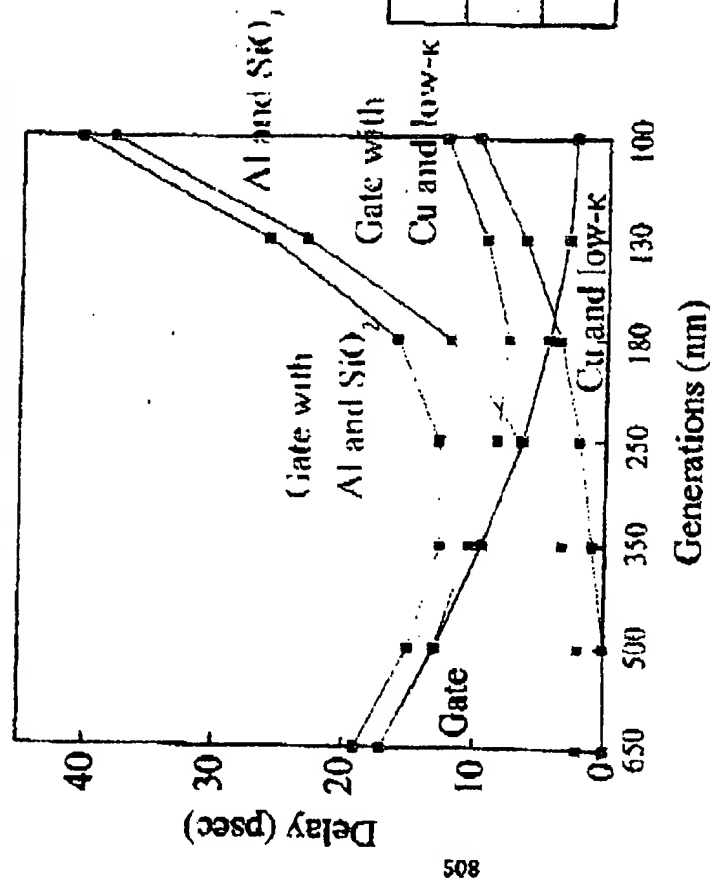
Exhibit I

Proceedings From The Ninth Meeting of
**THE SYMPOSIUM ON POLYMERS
FOR MICROELECTRONICS**



Bantec Museum & Gardens
Warrington, Delaware

Why Low-K Dielectric Materials ?



1997 The National Technology Roadmap for Semiconductors

| | 1999 | 2003 | 2009 |
|----|-------|-------|------|
| nm | 180 | 130 | 70 |
| K | 2.5-3 | 1.5-2 | ≤1.5 |

- Reduced RC time delay and cross-talking
 - Copper metal lines ($\rho_{Cu}=1.673 \mu\Omega\cdot\text{cm}$ vs. $\rho_{Al}=2.655 \mu\Omega\cdot\text{cm}$)
 - Low κ materials ($\kappa < 3$ vs. $\kappa_{SiO_2} \sim 4.0$)
 - New interconnect architecture (damascene/dual-damascene)

LOW-k DIELECTRIC MATERIALS

| Material | Dielectric Constant | Application Technique |
|---------------------------------|---------------------|-----------------------|
| Silicon dioxide | 3.9 - 4.1 | VD |
| Fluorinated oxide | 3.4 - 4.1 | VD |
| Polyimides | 3.0 - 3.6 | SO |
| Spin-on-glasses | 2.7 - 3.1 | SO |
| Fluorinated polyimides | 2.6 - 2.9 | SO |
| DLC (diamond-like carbon) | 2.8 - 3.0 | VD |
| Poly(arylene ethers) | 2.6 - 2.9 | SO |
| F-DLC | 2.4 - 2.9 | VD |
| Poly(arylenes) | 2.6 - 2.8 | SO |
| Poly(benzocyclobutene) | 2.6 - 2.8 | SO |
| Parylene N | 2.6 - 2.8 | VD |
| Poly(norbornenes) | 2.5 - 2.7 | SO |
| Polyimide-SSQ hybrids | 2.7 - 3.0 | SO |
| Parylene-AF ₄ | 2.3 - 2.5 | VD |
| Alkyl silanes/N ₂ O | 2.4 - 2.7 | VD |
| Ultra Low k Materials (k ≥ 2.2) | | |
| Teflon-AF | 1.9 - 2.1 | SO |
| Teflon-microemulsion | 1.9 - 2.1 | SO |
| Porous Materials | | |
| Polyimide nanofoams | 2.2 | SO |
| Silica aerogels | 1.1 - 2.2 | SO |
| Silica xerogels | 1.5 - 2.2 | SO |
| Mesoporous silica | 1.9 - 2.2 | SO |

Exhibit J

Dr. Stanley Wolf



PROCESS INTEGRATION FOR SUBMICRON IC TECHNOLOGIES

***Contract Course
Chartered Semiconductor - Singapore***

***Instructor:
Dr. Stanley Wolf***

***UC Berkeley Extension
Summer 1995***

Overview of Planarization Technologies

SOG types and properties:

- Silicate (inorganic)
- Phosphosilicate (inorganic)
- Siloxane (organic)
- Phosphosiloxane (organic)

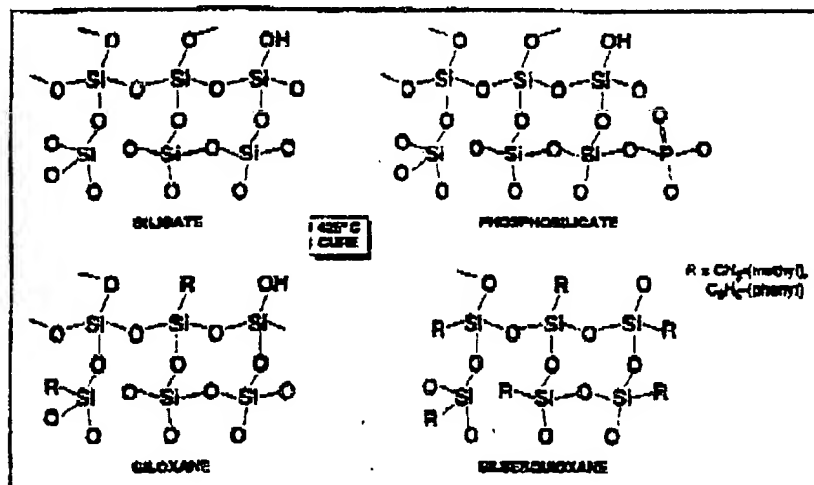


Figure 34. Chemical structure of various SOG material.

Inorganic SOG:

- Excellent resistance to oxygen plasmas
- Very high shrinkage on curing (doped with carbon to reduce that)

Organic SOG:

- Cannot withstand oxygen plasmas
- Low shrinkage characteristics

In short:

- Inorganic SOG's can't exceed 3000 Å in thickness
- Organic SOG's can't be exposed to oxygen plasma, and has a poor high temperature stability.

Exhibit K

**SILICON PROCESSING
FOR
THE VLSI ERA**

**VOLUME 1:
PROCESS TECHNOLOGY
Second Edition**

**STANLEY WOLF Ph.D.
RICHARD N. TAUBER Ph.D.**

**LATTICE PRESS
Sunset Beach, California**

792 SILICON PROCESSING FOR THE VLSI ERA

There are several processes that need to be performed on the dielectric films after their deposition. The films must be able to be properly integrated into such processing. First, the films will need to undergo CMP, and such materials will thus need to be polishing-process-compatible. Next, vias (and perhaps grooves in a damascene technology) will have to be etched in the films. Issues of etch chemistry, etch selectivity to oxides, nitrides, and oxynitrides will need to be addressed. After etching, the resist mask must be removed. It must be possible to perform any resist-removal processing (and any resist-residue-removal steps) without damaging the dielectric layers.

Other *miscellaneous process integration* issues of low- k dielectrics include the following. First, although the gap-fill characteristics of the low- k films are not a consideration for damascene structures (as the dielectric films are deposited on surfaces that have been planarized by CMP), they will need to be deposited in a single pass at a thickness of about twice that used in dielectric films in non-damascene structures. Thus, they will need to exhibit crack-resistant characteristics for such film thicknesses. In addition, for some dielectric materials it might be necessary to use a capping layer of conventional CVD oxide or oxynitride to make the overall layer able to withstand the CMP process. For non-damascene structures (i.e., in which the metal [Al] patterns are etched and then covered with a dielectric layer), the gap-fill characteristics of the low- k material will be important. It is asserted that both damascene and non-damascene interconnect structures will continue to find use as devices are scaled, depending on the circuit application. Finally, use of spin-on dielectric films that evolve toxic vapors will be avoided.

15.10.2 First Generation Low- k Dielectrics ($2.8 < k < 3.5$)

Dielectric materials with k values that are moderately lower than SiO_2 were the first materials to be adopted for this function. Three types of such moderately low- k films, *HSQ films* (introduced in Sect. 15-9), *fluorinated oxide films** and *low- k SOGs* found their way into IC production by 1990. Their main use was reported to be in the $0.5\ \mu\text{m}$ to $0.25\ \mu\text{m}$ CMOS generations.

The *spin-on HSQ films* were initially investigated as replacement films for siloxane SOG films in non-etchback SOG planarization processes. But they also exhibit a lower k -value than SiO_2 films (HSQ films have a k value of < 2.9). The HSQ material is available from Dow Corning under the tradename of FOx (Flowable Oxide).¹³⁷ HSQ was reported to have been integrated into products by Texas Instruments in the $0.5\ \mu\text{m}$ generation, and in 1997 an article describing its use in a five-layer Al-interconnect structure was published.¹¹⁸ An oxide cap was used on the HSQ film to improve its low- k value stability during W-CVD and Al fill.

FSG films are silicon oxyfluorides (F_xSiO_y) deposited by CVD. They exhibit a k value of ~ 3.5 . FSG films are formed in PECVD or HDP-CVD tools by adding SiF_4 to the process gas ambient used to deposit SiO_2 by CVD (silane and oxygen). By increasing the SiF_4 flowrate, more F is incorporated into the film. Higher concentrations of F cause the value of k to decrease. However, a maximum of about 6% F is permitted to be incorporated into the FSG films because higher concentrations would cause F to be evolved during RIE etch of these films. This would corrode the metal in the interconnect structure. FSG films were reported to be successfully integrated into 0.35 and $0.25\ \mu\text{m}$ circuits by IBM.⁷⁵ The evolution of F from the FSG oxide has

* These are also referred to as fluorosilicated glass (FSG), or silicon oxyfluoride (SiOF).

MULTI-LEVEL INTERCONNECTS FOR ULSI 793

raised concerns about using this material in Cu interconnect systems. The fluoride attacks the Ta-based liners of Cu systems, leading to volatile TaF_5 formation, and the consequent loss of adhesion between the low- k film and the Ta. A low- k SOG film is available from Allied-Signal with a k value of 2.9. It is an inorganic siloxane polymer named Accuspin T-23. When cured at 380–400°C, it can tolerate temperatures as high as 450°C before decomposing.

15.10.3 Second-Generation Low- k Dielectrics ($2.5 < k < 2.8$)

The largest variety low- k dielectrics is found in the second generation of such substances, which exhibit k values between 2.5 and 2.8. It is believed that these materials will begin to be incorporated into ICs at the 0.18 μm generation. Both *spin-on* and *CVD* low- k materials have been developed within this range of permittivities. Both types are described here.

15.10.3.1 2nd-Gen Spin-On Dielectrics with $2.5 < k < 2.8$: Most spin-on organic low- k polymers¹¹⁷ are significantly different from their conventional SOG counterparts. That is, they do not evolve moisture during curing. Because they exhibit better crack resistance they can be applied in a single pass to a greater thickness. Three organic spin-on low- k dielectrics being considered are: *poly(arylene) ethers* (PAE), *fluoro-polyimides*, and *bis-benzocyclobutenes* (BCB). In addition, there is an organo-siloxane polymer type of film. Spin-on dielectric films are deposited without the use of an expensive CVD tool (they need a track cluster tool with spinners and hot plates instead). The cost of the spin-on liquid, however, is typically much higher per wafer than the cost of the CVD reactant gases.¹³⁸ In addition, a 100-nm-thick capping SiO_2 layer (deposited by CVD) will probably still be needed with spin-on low- k layers. Finally, spin-on films reduce the topology of any previous steps by more than 50%.

The PAE materials are aromatic structures which have k values of about 2.5 and also have the ability to withstand high temperatures. A fluorinated PAE product from Allied-Signal is named FLARE, and has a k value of ~2.8. It was designed to have low outgassing characteristics and high thermal and mechanical stability.¹¹⁹ Dow Chemical has a similar material named SiLK. A report by Fujitsu detailed their experience with integrating FLARE and SiLK into a two-level Cu damascene structure. They showed little change in k value after a five-hour anneal at 425°C.¹²⁰ Schumacher has pioneered the use of non-fluorinated PAE materials, with their latest such product being VELOX. This material can be deposited without an adhesion promoter and does not require a capping layer. A fluoro-polyimide named FPI-136M with $k = 2.7$ is sold by DuPont.

BCB is a co-polymer of divinyl-siloxane and bis-benzocyclobutene (which is often referred to as DVS-BCB, or simply BCB). It is a thermoset silicon-containing resin that exhibits a k value of 2.7, but which has limited thermal stability even at temperatures of 350°C. Nevertheless, there are no by-products to the polymerization reaction and no corrosive or volatile by-products are evolved during the cure. BCB demonstrates excellent gap-fill, adhesion, and planarization capabilities. Dow Chemical offers a BCB product for IC ILD applications called Cyclotene. A study in 1998 indicated that Cu drifts much more slowly through BCB films than through PAE.¹²¹ Thus, use of BCB films could relax the requirements of metal diffusion barriers. But its limited thermal stability constrains its use to applications with a reduced thermal ceiling.

The SOG-like material in this class is Accuspin T-24 from Allied-Signal. It is a hydrido-organosiloxane polymer (HOSP) which has a k value of 2.5 after curing. This HOSP has a 12%

Exhibit 2

A STUDY OF PLASMA TREATMENTS ON SILOXANE SOG

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ABSTRACT

In this paper, the mechanisms of siloxane SOG damaged by oxygen and other plasma treatments are studied with Fourier transformation infrared spectrum (FTIR), secondary ion mass spectrum (SIMS), refractive index (R.I.), and the amount of SOG loss. The information of water absorption is provided with residual gas analysis (RGA). The relationship between the extent of SOG damaged by plasma treatment with moisture absorption and SOG loss are investigated. Oxygen plasma treatment oxidizes siloxane SOG and results in high moisture absorption. Several approaches have been tried to reduce the moisture absorption caused by oxygen plasma treatment. Ar plasma treatment is found to be the most effective one to prevent the degradation of SOG by oxygen plasma. After implementing Ar plasma treatment into processing, lower and stable via contact resistance can be achieved; good yield has also been demonstrated.

INTRODUCTION

In the fabrication of sub-micron devices, the planarization of inter-metal dielectric (IMD) films have become increasingly important. Among various planarization schemes, the spin-on-glass (SOG) process is a very popular approach [1-3]. Due to the gap filling capability and planarization performance, siloxane SOG is the major material used in ILD scheme for sub-micron devices. However, there are many concerns of siloxane SOG. One of them is moisture absorption, especially after exposed to oxygen plasma [4,5]. During the stripping of via photo resist, the SOG at via side wall is exposed to oxygen plasma. Therefore, poisoned via and SOG retrogression are potential issues that may impact chip yield or reliability [6-8]. One of the approaches has to convert the surface layer of organic SOG into inorganic type by N₂O or O₂ plasma treatment and followed by a outgassing process [5,9].

In this study, we investigated the effect of plasma treatments on siloxane SOG in order to find out approaches to strengthen SOG from damages by oxygen plasma. Plasma treatment by Ar is found most effective to avoid SOG degradation by oxygen plasma, and can result in good electrical and reliability performances.

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EXPERIMENTAL

There are two kinds of methyl polysiloxane SOG used in this study: Allied Signal Accuglass 111 type and Accuglass 314. Various plasma treatments on cured SOG had been studied. The plasma treatments of N₂ or N₂O was performed in the CVD tool with process temperature of 400C. The O₂ plasma treatments were performed in two different systems: a batch type asher with high power and a single wafer asher with low power. The Ar plasma treatment was conducted in sputtering chamber.

The residual gas analyzer (RGA) was used to detect the outgassing of SOG films. The composition profiles of carbon, hydrogen and oxygen inside SOG were analyzed with secondary ion mass spectrum (SIMS). Fourier transformation infrared spectrum (FTIR) was taken to provide the informations of bonding structure and moisture absorption of SOG films. Both Prometric and Ellipsometer were used to measure the film thickness. The refractive index (R.I.) was measured with Ellipsometer. In the evaluation of the electrical and reliability performances, devices with deep via structure were used.

RESULTS AND DISCUSSION

Firstly, the time dependence of moisture absorption of cured SOG was studied in order to set a proper queue time control between SOG curing and the deposition of oxide capping layer. The water content inside cured SOG films of different exposure time was quantified by comparing the area ratio of Si-OH to Si-O peaks from FTIR measurement. As shown in Figure 1, the SOG films cured at 400, 425 or 450C all have similar trend of moisture absorption: fast absorption at the initial several hours after curing and getting saturation after 24 hours. Considering the fast absorption, the queue time control is not a really good approach for manufacturing. As shown in the RGA data in Figure 2, the amount of H₂O in SOG is found related to the film thickness of SOG, indicating that the moisture absorption is through the bulk SOG not only on the SOG surface.

The moisture absorption of SOG damaged by O₂ plasma treatment can be several orders higher than that of cured SOG, as shown in Figure 2. Other than serious moisture absorption, O₂ plasma treatment may also be responsible for the retrogression of SOG (Figure 3) at the sidewall of via holes. Unfortunately, the impact of oxygen plasma on SOG is almost unavoidable if SOG is exposed at via sidewall; ashing treatment is commonly applied in stripping the photo resist after via etching. The wet cleaning with D.I. water rinse is usually applied after the dry ashing. The SOG damaged by O₂ plasma can absorb higher amount of moisture than the normal SOG, and may result in greater extent of outgassing at sequential process. Either the retrogression or outgassing of SOG films has higher potential in degrading chip yield and reliability. Therefore efforts had been put on to understand the degradation mechanism of SOG by O₂ plasma and to find a method to reduce the harmful effects of O₂ plasma damages, especially moisture absorption.

With FTIR and SIMS, the O₂ plasma treatment is found to result in the oxidation of the organic group of siloxane SOG. In other words, the siloxane SOG is converted to inorganic SOG. As shown in the FTIR spectra in Figure 4, the Si-CH₃ peak in siloxane SOG disappears after O₂ plasma treatment and the Si-O-Si peak

Exhibit M

AlliedSignal Spin-on Products

Comparison of Film Properties

| Product | Material | Thickness (μm) | Thickness (Å) | Refractive Index | Modulus (GPa) | Adhesion (MPa) | Reliability (h) | Primary Application | |
|---------|----------------------|----------------|---------------|------------------|---------------|----------------|-----------------|-------------------------------------|---------------------------------------|
| USG-50 | Silicate | 6-7 | 2150Å | N/A | | Fair | Very High | Poly Smoothing | |
| 103AS | Silicate | 6-8 | 1370Å | 12-14% | | Fair | Very High | Poly Smoothing | |
| 803AS | Silicate | 6-7 | 2450Å | 12-14% | | Fair | Very High | Poly Smoothing | |
| P-062A | Phosphosilicate | 4.1 | 645Å | 16-18% | ≥0.5 | Fair | High | Poly/Metal Planarization | |
| P-082A | Phosphosilicate | 4.1 | 820Å | 16-18% | ≥0.5 | Fair | High | Poly/Metal Planarization | |
| P-112A | Phosphosilicate | 4.1 | 1025Å | 16-18% | ≥0.5 | Fair | High | Poly/Metal Planarization | |
| P-064A | Phosphosilicate | 4.3 | 600Å | 16-18% | ≥0.5 | Fair | High | Poly/Metal Planarization | |
| P-094A | Phosphosilicate | 4.3 | 800Å | 16-18% | ≥0.5 | Fair | High | Poly/Metal Planarization | |
| P-114A | Phosphosilicate | 4.3 | 1080Å | 16-18% | ≥0.5 | Fair | High | Poly/Metal Planarization | |
| 111 | Methylsiloxane | 3.8 | 1285Å | 4-6% | ≥0.3 | Good | Medium | Metal Partial Etchback/Non-Etchback | |
| 211 | Methylsiloxane | 3.8 | 2170Å | 4-6% | ≥0.3 | Good | Medium | Metal Partial Etchback/Non-Etchback | |
| 311 | Methylsiloxane | 3.8 | 3140Å | 4-6% | ≥0.3 | Good | Medium | Metal Partial Etchback/Non-Etchback | |
| 512 | Methylsiloxane | 3.1 | 5000Å | 2% | ≥0.3 | Good | Low | Metal Partial Etchback/Non-Etchback | |
| 512B | Methylsiloxane | 3.1 | 5100Å | 2% | ≥0.3 | Good | Low | Metal Partial Etchback/Non-Etchback | |
| 214 | Methylsiloxane | 3.8 | 2120Å | 13% | ≥0.15 | Very Good | Medium | Metal Partial Etchback/Non-Etchback | |
| 314 | Methylsiloxane | 3.8 | 2865Å | 13% | ≥0.15 | Very Good | Medium | Metal Partial Etchback/Non-Etchback | |
| 218 | Methyltrisiloxane | 2.7 | 2000Å | 2% | ≥0.1 | Excellent | Fair | Very Low | Metal Partial Etchback |
| 418 | Methyltrisiloxane | 2.7 | 4000Å | 2% | ≥0.1 | Excellent | Fair | Very Low | Metal Partial Etchback |
| 720 | Methylphenylsiloxane | 3.0 | 2700Å | 3% | ≥0.1 | Excellent | | Very Low | Metal Partial Etchback/Total Etchback |
| 1513EL | Organic Polymer | N/A | 15000Å | N/A | | Excellent | N/A | Regional Total Etchback | |
| 913EL | Organic Polymer | N/A | 9000Å | N/A | | Excellent | N/A | Regional Total Etchback | |
| 713EL | Organic Polymer | N/A | 6800Å | N/A | | Excellent | N/A | Regional Total Etchback | |
| 412EL | Organic Polymer | N/A | 4000Å | N/A | | Excellent | N/A | Regional Total Etchback | |



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EXHIBIT N

FLUORINE-IMPLANTED TREATMENT (FIT) SOG FOR THE NON-ETCHBACK INTERMETAL DIELECTRIC

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ABSTRACT

In this work we describe a new fluorine-implanted treatment (FIT) SOG modification technology to obtain the superior fluorinated SiO_2 film. The properties of FIT SOG were examined by measuring the index of refraction, thickness shrinkage, temperature dependence of stress and wet etching rate. The outgassing behavior of FIT SOG was determined by means of residual gas analyzer (RGA) method for the application of non-etchback SOG process. We have applied the FIT SOG for the 16M DRAM non-etchback intermetal dielectric (IMD) planarization process, and excellent electric performances were obtained.

INTRODUCTION

Spin-On-Glass (SOG) films are often used in the intermetal dielectric between the first level aluminum interconnection and the second one for the purpose of planarization in the ULSI multilevel interconnection fabrication. The most commonly used process for SOG planarization is etch back process that involves depositing a thin CVD oxide upon the aluminum metal, spun the SOG and then etching away the film only leaves the SOG to fill the metal trench. Then an encapsulated CVD oxide is deposited to complete the IMD planarization step. While the etch back process work well in current IMD planarization technology, the non-etch back processing is highly desirable because of its greater simplicity and wider process window. The most concern for non-etchback processing is the issues of "poisoned via". Poisoned via is a contaminated area resulting from SOG residues that cause poor contact between metal 1 and metal 2 interconnect. Several investigations have shown the "poisoned via" was attributed to the presence of moisture that was either formed within the SOG due to insufficient cure or picked up from preclean steps, especially for the siloxane SOGs, as they contain much residual OH and much moisture adsorption. Recently, some authors proposed room temperature CVD technique [1] and SOG film modification processing [2] by using the fluorotriethoxysilane and fluoralkoxysilane as the gas source and coating catalyst. The purpose of these modifications is to obtain a fluorinated SiO_2 film. A fluorinated SiO_2 film shows superior dielectric properties compare to the conventional oxide. This paper proposes a new

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fluorine-implanted treatment (FIT) SOG modification technology to obtain the superior fluorinated SiO_2 film. The properties of FIT SOG were examined and compared to the conventional SOG films. The applications of FIT SOG for the non-etchback IMD planarization were described. Low via RC and 100% yield indicate there have no "poisoned via" issues by using FIT non-etchback SOG process.

EXPERIMENTAL DETAILS

To characterize the film's properties, the siloxane based SOG coating upon the 8 inch (100) Si wafer was adopted. The SOG was dispensed and baked on a commercial coating equipment. Each spin gives a thickness of approximately 2200Å when measured on a bare Si test wafer. Following bake, the wafers were carried out ion implantation step. Fluorine (F^+) was employed as the doping source in the ion implantation process. The acceleration voltage was 40 or 80 keV and the implanted doses were $3\text{--}5 \times 10^{15}$ atom/ cm^2 . An non-implanted type SOG wafers were prepared for the aim of measurement comparison. Finally, all the wafers (implanted and non-implanted) were cured at a maximum temperature of 425°C. Film thickness and index of refraction was measured by using an ellipsometer. The measurement of stress as a function of annealing temperature was done in a Tencor stress measurement equipment that employs a laser scanning system for measuring the radius of curvature of a wafer. Moisture desorption was studied by using of residual gas analyzer (RGA) thermal analysis [3] for the purpose of the application of non-etchback SOG process. The process flow of FIT non-etchback SOG process is listed in Table 1.

RESULTS AND DISCUSSION

The index of refraction has the value of 1.44 for the FIT SOG compare to the value of 1.40 for the non-modified SOG. Higher index of refraction indicates the modified film is denser than the control one with the assistance of fluorine implantation. This result coincides to the measurement of film shrinkage. The shrinkage of FIT SOG has the value of 20% and much larger than the value of 8% for the conventional SOG after curing process. Measurement of wet etching rate was carried out in 200:1 BOE chemistry. As shown in Fig. 1, the wet etching rate of FIT SOG is much lower than the non-modified SOG and even lower than the PECVD oxide and approximate to the value of thermal oxide. In summary the measurement of refractive index, thickness shrinkage and wet etching rate, we concluded that the FIT SOG is much denser than the conventional siloxane SOG. The typical temperature cycle stress variation of as-coated SOG is shown in Fig. 2. At about 250°C, the tensile stress increased sharply as moisture was evolved and finally reached a maximum of over 1×10^9 dyne/ cm^2 . With the moisture removed, the cooling behavior was found to be elastic. The thermal cycle was repeated after storing the wafer for several days in an air room environment. The stress measurement of FIT SOG is shown in Fig. 3. By comparison of Fig. 2 and Fig. 3, the FIT SOG behavior two different stress performances (1) compressive stress (2) without moisture desorption stress variation that often observed in siloxane SOG as mentioned before (3) little stress increase during thermal cycle. The temperature-stress measurement was carried out after the film is allowed to stand in air room for 10 days' period and O_2 plasma treatment and is shown in Fig. 4 and Fig. 5, individually. The stress measurement suggests the film scarcely adsorbs the moisture even after O_2 plasma process. This argument was examined by using the

RGA thermal analysis. As shown in Fig. 6, the amount of moisture desorption of FIT SOG is purely insignificant compare to that of non-modified SOG. The application results of FIT SOG to interlayer dielectric film application of Al interconnect in our 16M DRAM product, without using any SOG etchback process are shown in Fig. 7 and Fig. 8. Low via RC and 100% yield were obtain. In general, the RC value of FIT non-etchback process is lower than that of etchback process, especially for the 0.5 μm via size. That suggests there has no "poisoned via" issues by using FIT non-etchback SOG for the application of intermetal dielectric planarization.

CONCLUSIONS

A fluorine-implanted treatment (FIT) SOG modification technology was proposed for the application of intermetal dielectric planarization. The modified FIT SOG has the following character: (1) more dense the film compares to SOG (2) very low wet etching rate (3) compressive stress behavior (4) little stress increase during temperature treatment (5) less moisture adsorption. The application results of FIT SOG to interlayer dielectric film application of Al interconnect shows the FIT SOG is superior to the non-etchback process for the application of intermetal dielectric planarization.

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Table 1 The process flow of FIT non-etchback intermetal dielectric planarization.

| Process Step | Specification or Process Parameter |
|--------------------------------|---|
| 1. Metal 1 sputter | 6000 Å, Al plug |
| 2. PE TEOS CVD | 2000 Å |
| 3. SOG spin and bake | 4000 Å, double coating |
| 4. F ⁺ implantation | 40 or 80 Kev, 3E15 atom/cm ² |
| 5. Curing in furnace | 425 °C, 30 minute |
| 6. PE TEOS CVD | 4500 Å |
| 7. Metal 2 sputtering | 8000 Å, AlSiCu |